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OM protein - protein search, using sw model

Run on: July 27, 2004, 10:02:56 ; Search time 54 Seconds  
(without alignments)  
1104.028 Million cell updates/sec

Title: US-09-813-820-4  
Perfect score: 1122  
Sequence: 1 MRGSHHHHGGSDKVATIT.....GIEGTVKGLKVLKQDKDK 211

Scoring table: BLOSUM62  
Gapop 10.0 , Gapext 0.5

Searched: 1586107 seqs, 282547505 residues

Total number of hits satisfying chosen parameters: 1586107

Minimum DB seq length: 0  
Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%  
Maximum Match 100%  
Listing first 45 summaries

Database : A\_Geneseq\_29Jan04:\*  
1: Geneseqp1980s:\*  
2: Geneseqp1990s:\*  
3: Geneseqp2000s:\*  
4: Geneseqp2001s:\*  
5: Geneseqp2002s:\*  
6: Geneseqp2003as:\*  
7: Geneseqp2003bs:\*  
8: Geneseqp2004s:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	1122	100.0	211	2 AAW31553	Aaw31553 Collagen
2	1054.5	94.0	512	2 AAW31554	Aaw31554 Collagen
3	1048	93.4	1183	6 ABU79084	Abu79084 S. aureus
4	904	80.6	1185	2 AAR22675	Aar22675 Collagen
5	884	78.8	168	4 AAE11855	Aae11855 Staphyloc
6	830	74.0	159	2 AAW31552	Aaw31552 Collagen
7	491	43.8	731	7 ADC97372	Adc97372 E. faeciu
8	178	15.9	458	3 AAB37667	Aab37667 Collagen
9	178	15.9	458	7 ADD90648	Add90648 Enterococ
10	165.5	14.8	146	3 AAB37668	Aab37668 Collagen
11	154	13.7	168	4 AAE11856	Aae11856 Enterococ
12	141.5	12.6	316	7 ABM79017	Abm79017 Staphyloc
13	131.5	11.7	343	7 ABM79016	Abm79016 Staphyloc
14	125.5	11.2	345	2 AAW31555	Aaw31555 Fibronect
15	119	10.6	336	5 AAB53995	Abb53995 Lactococc
16	118	10.5	560	7 ABM79015	Abm79015 Staphyloc
17	116	10.3	473	2 AAW68400	Aaw68400 Clostridi
18	114.5	10.2	688	2 AAY00241	Aay00241 Enterococ
19	114.5	10.2	688	5 ABP43460	Abp43460 E faecali
20	114.5	10.2	688	6 ABU88488	Abu88488 E. faecal
21	114.5	10.2	688	6 ABU13739	Abu13739 Enterococ
22	114.5	10.2	2032	2 AAY00238	Aay00238 Enterococ
23	114.5	10.2	2032	2 AAY00240	Aay00240 Enterococ
24	114.5	10.2	2032	2 AAY00242	Aay00242 Enterococ
25	114.5	10.2	2032	5 ABP43459	Abp43459 E faecali

26	114.5	10.2	2032	5 ABP43457	Abp43457 E faecali
27	114.5	10.2	2032	5 ABP43461	Abp43461 E faecali
28	114.5	10.2	2032	6 ABU88489	Abu88489 E. faecal
29	114.5	10.2	2032	6 ABU88485	Abu88485 E. faecal
30	114.5	10.2	2032	6 ABU88487	Abu88487 E. faecal
31	114.5	10.2	2032	6 ABU13740	Abu13740 Enterococ
32	114.5	10.2	2032	6 ABU13736	Abu13736 Enterococ
33	114.5	10.2	2032	6 ABU13738	Abu13738 Enterococ
34	114	10.2	200	6 ABP58754	Abp58754 Recombina
35	113.5	10.1	184	2 AAY32917	Aay32917 E.coli op
36	113.5	10.1	184	3 AAB10309	Aab10309 Human KGF
37	113.5	10.1	184	4 AAB61673	Aab61673 PQ560-Cys
38	113.5	10.1	184	6 ABP54289	Abp54289 PQ560-Cys
39	113.5	10.1	184	6 ADA95466	Ada95466 Synthetic
40	113.5	10.1	184	7 ADD66139	Add66139 Human mat
41	113.5	10.1	1788	6 ABU18740	Abu18740 Protein e
42	113	10.1	303	5 ABG67279	Abg67279 Polythoa
43	111.5	9.9	2627	6 ABR39483	Abr39483 L. cuprin
44	111.5	9.9	3333	6 ABR39482	Abr39482 L. cuprin
45	110.5	9.8	1338	2 AAR41731	Aar41731 High mole

ALIGNMENTS

RESULT 1

AAW31553	ID	AAW31553 standard; protein; 211 AA.
XX	AC	AAW31553;
XX	DT	27-AUG-2003 (revised)
DT	25-MAR-2003	(revised)
DT	21-MAY-1998	(first entry)
XX	DE	Collagen binding protein M31 epitope.
XX	KW	Collagen binding protein; cna gene; sepsis; infection;
KW	KW	microbial surface component recognising adhesive matrix molecule; MSCRAMM;
KW	KW	adhesin; vaccine; immunisation; diagnosis; therapy; epitope M31.
XX	OS	Staphylococcus aureus.
XX	Key	Location/Qualifiers
FT	Peptide	1..12
FT	Protein	/note= "vector PQE30-derived peptide"
FT	Protein	13..211
FT	Protein	/note= "epitope M31"
XX	XX	WO9743314-A2.
XX	XX	20-NOV-1997.
XX	XX	14-MAY-1997; 97WO-US008210.
XX	XX	16-MAY-1996; 96US-0017678P.
XX	XX	(TEXA ) UNIV TEXAS A & M SYSTEM.
XX	XX	(UABR-) UAB RES FOUND.
XX	XX	Hoeoek M, Patti JM, House-Pompeo K, Sthanam N, Symersky J;
XX	XX	WPI; 1998-008801/01.
XX	XX	N-PSDB; AAT93437.
XX	XX	Antibody that interacts with collagen binding domain of Staphylococcal
XX	XX	cna gene product - useful to prevent bacterial sepsis in animal infected
XX	XX	with Staphylococcus aureus.
XX	XX	Claim 31; Page 115-116; 143pp; English.
XX	XX	This protein comprises Staphylococcus aureus collagen binding protein
XX	XX	(CBP) epitope M31, i.e. amino acids 61-343 of full-length CBP, plus a

CC vector-derived N-terminal peptide. Claimed 441, 849 and 1500 bp nucleic  
 CC acid sequences (see AAT93436-38) respectively encode CBP epitopes M17,  
 CC M31 and M55 (see AAW31552-54) that confer protection against *S. aureus*  
 CC infection. These nucleic acid sequences can be used in the recombinant  
 CC production of the CBP epitopes. The CBP protein and antigenic epitopes  
 CC are contemplated for use in the treatment of pathological infections,  
 CC especially to prevent bacterial adhesion to collagen. The claimed nucleic  
 CC acids as well as claimed anti-CBP antibodies will also be of use in  
 CC screening, diagnostic and therapeutic applications including active and  
 CC passive immunisation and methods for the prevention of bacterial  
 CC colonisation in an animal such as a human. The CBP epitopes are also  
 CC contemplated for use in the preparation of vaccines and as carrier  
 CC proteins in vaccine formulations, as well as in the formulation of  
 CC compositions for the prevention of *S. aureus* infection. (Updated on 25-  
 CC MAR-2003 to correct PI field.) (Updated on 27-AUG-2003 to correct OS  
 CC field.)  
 XX Sequence 211 AA;  
 SQ

Query Match 100.0%; Score 1122; DB 2; Length 211;  
 Best Local Similarity 100.0%; Pred. No. 9.2e-86;  
 Matches 211; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
 1 MRGSHHHHGGSDKVAITTSNGKSTNTVHKSEAGTSVFFYKTDMLPDTTHVRWFL 60  
 1 MRGSHHHHGGSDKVAITTSNGKSTNTVHKSEAGTSVFFYKTDMLPDTTHVRWFL 60  
 61 NINNEKSYVSKDITIKDQIQGGQQLDLSTLINVGTGHSNYSGQSATDPEKAFPGSKI 120  
 61 NINNEKSYVSKDITIKDQIQGGQQLDLSTLINVGTGHSNYSGQSATDPEKAFPGSKI 120  
 121 TVDNTKNTIDVTIPQGYGSYNSFSINYKTKITNEQKEFVNNSQAWYQEHGKEEVNGKSF 180  
 121 TVDNTKNTIDVTIPQGYGSYNSFSINYKTKITNEQKEFVNNSQAWYQEHGKEEVNGKSF 180  
 181 NHTVHNINANAGIEGTGKELKVLKQDKDTK 211  
 181 NHTVHNINANAGIEGTGKELKVLKQDKDTK 211

RESULT 2  
 AAW31554  
 ID AAW31554 standard; protein; 512 AA.  
 AC AAW31554;  
 XX  
 XX 27-AUG-2003 (revised)  
 DT 25-MAR-2003 (revised)  
 DT 21-MAY-1998 (first entry)  
 XX  
 XX Collagen binding protein M55 epitope.  
 DE  
 XX Collagen binding protein; cna gene; sepsis; infection;  
 KW microbial surface component recognising adhesive matrix molecule; MSCRAMM;  
 KW adhesin; vaccine; immunisation; diagnosis; therapy; epitope M55.  
 XX  
 XX Staphylococcus aureus.  
 OS  
 XX  
 XX Key Location/Qualifiers  
 FH Peptide 1..12  
 FT /note= "vector pQE30-derived peptide"  
 FT 13..512  
 FT /note= "epitope M17"  
 XX  
 XX WO9743314-A2.  
 XX  
 XX 20-NOV-1997.  
 FD  
 XX 14-MAY-1997; 97WO-US008210.  
 XX  
 XX 16-MAY-1996; 96US-0017678P.  
 XX  
 XX (TEXA ) UNIV TEXAS A & M SYSTEM.  
 PA

PA (UABR-) UAB RES FOUND.  
 XX  
 XX Hoeoek M, Patti JM, House-Pompeo K, Sthanam N, Symersky J;  
 XX WPI: 1998-008801/01.  
 DR N-PSDB; AAT93438.  
 XX  
 XX Antibody that interacts with collagen binding domain of Staphylococcal  
 PT cna gene product - useful to prevent bacterial sepsis in animal infected  
 PT with Staphylococcus aureus.  
 XX  
 PS Claim 31; Page 117-119; 143pp; English.  
 XX  
 CC This protein comprises Staphylococcus aureus collagen binding protein  
 CC (CBP) epitope M55, i.e. amino acids 30-531 of full-length CBP, plus a  
 CC vector-derived N-terminal peptide. Claimed 441, 849 and 1500 bp nucleic  
 CC acid sequences (see AAT93436-38) respectively encode CBP epitopes M17,  
 CC M31 and M55 (see AAW31552-54) that confer protection against *S. aureus*  
 CC infection. These nucleic acid sequences can be used in the recombinant  
 CC production of the CBP epitopes. The CBP protein and antigenic epitopes  
 CC are contemplated for use in the treatment of pathological infections,  
 CC especially to prevent bacterial adhesion to collagen. The claimed nucleic  
 CC acids as well as claimed anti-CBP antibodies will also be of use in  
 CC screening, diagnostic and therapeutic applications including active and  
 CC passive immunisation and methods for the prevention of bacterial  
 CC colonisation in an animal such as a human. The CBP epitopes are also  
 CC contemplated for use in the preparation of vaccines and as carrier  
 CC proteins in vaccine formulations, as well as in the formulation of  
 CC compositions for the prevention of *S. aureus* infection. (Updated on 25-  
 CC MAR-2003 to correct PI field.) (Updated on 27-AUG-2003 to correct OS  
 CC field.)  
 XX  
 SQ Sequence 512 AA;  
 Query Match 94.0%; Score 1054.5; DB 2; Length 512;  
 Best Local Similarity 64.7%; Pred. No. 1.3e-79;  
 Matches 211; Conservative 0; Mismatches 0; Indels 115; Gaps 1;  
 1 MRGSHHHHGGSDKVAITTSNGKSTNTVHKSEAGTSVFFYKTDMLPDTTHVRWFL 60  
 1 MRGSHHHHGGSDKVAITTSNGKSTNTVHKSEAGTSVFFYKTDMLPDTTHVRWFL 60  
 121 TVDNTKNTIDVTIPQGYGSYNSFSINYKTKITNEQKEFVNNSQAWYQEHGKEEVNGKSF 180  
 121 TVDNTKNTIDVTIPQGYGSYNSFSINYKTKITNEQKEFVNNSQAWYQEHGKEEVNGKSF 180  
 66 KSYVSKDITIKDQIQGGQQLDLSTLINVGTGHSNYSGQSATDPEKAFPGSKI 125  
 181 KSYVSKDITIKDQIQGGQQLDLSTLINVGTGHSNYSGQSATDPEKAFPGSKI 240  
 126 KNTIDVTIPQGYGSYNSFSINYKTKITNEQKEFVNNSQAWYQEHGKEEVNGKSF 185  
 241 KNTIDVTIPQGYGSYNSFSINYKTKITNEQKEFVNNSQAWYQEHGKEEVNGKSF 300  
 186 NINANAGIEGTGKELKVLKQDKDTK 211  
 301 NINANAGIEGTGKELKVLKQDKDTK 326

RESULT 3  
 ABU79084  
 ID ABU79084 standard; protein; 1183 AA.  
 XX  
 XX AC ABU79084;  
 XX  
 XX 18-JUN-2003 (first entry)  
 DT  
 XX  
 XX S. aureus collagen adhesin protein.  
 DE  
 XX

KW Superantigen; SAg; staphylococcal enterotoxin; tumour; cancer; apoptosis;  
KW gene therapy; mammalian cell receptor; cytostatic;  
KW tumour associated lipid; energy; T cell; antigen presenting cell; APC;  
KW tumouricidal immunocyte; antitumour.  
XX  
XX  
OS Staphylococcus aureus.  
XX  
PN US2002177551-A1.  
XX  
XX N-PSDB; ACA64715.  
PD 28-NOV-2002.  
XX  
XX 30-MAY-2001; 2001US-00870759.  
XX  
XX 31-MAY-2000; 2000US-0208128P.  
PR (TERM/) TERMAN D S.  
XX  
XX Terman DS;  
XX  
XX WPI; 2003-361759/34.  
DR N-PSDB; ACA64715.  
XX  
XX A mammalian cell receptor, useful in the treatment of cancer by binding  
PT to tumor associated lipids where the binding induces energy or apoptosis  
PT in T cells and antigen presenting cells.  
XX  
XX Disclosure; Page; 167pp; English.  
XX  
XX The invention relates to a mammalian cell receptor, useful in the  
CC treatment of cancer, which binds to tumour associated lipids and induces  
CC energy or apoptosis in the T cells and antigen presenting cells (APCs).  
CC Also included are a mammalian cell useful in the treatment of cancer  
CC where the receptor which binds tumour associated lipids and induces  
CC cellular inactivation or death is deleted or functionally deactivated,  
CC producing (M1) a tumouricidal immunocyte population in vivo in a mammal  
CC (by allowing tumour associated lipids to contact immunocytes in which  
CC receptors for immunosuppressive fatty acids, ceramides, glycolipids,  
CC sphingolipids, glycosphingolipids, phosphoglycolipids, gangliosides,  
CC sialylated glycans, lipopeptides and proteoglycolipids are inactivated or  
CC deleted), a construct useful in the treatment of cancer comprising a  
CC superantigen (SAg) nucleotide inserted into a virus, a mammalian T cell  
CC useful in the treatment of cancer (where an adaptor protein which  
CC inhibits T cell activation by tumour associated antigens is deleted or  
CC functionally deactivated), a composition useful in the treatment of  
CC cancer (comprising a lipid raft conjugated to a superantigen), producing  
CC (M2) a tumouricidal immunocyte population ex vivo in a mammal (by  
CC allowing tumour associated lipids to contact immunocytes, in which  
CC receptors for the lipids are inactivated or deleted to produce a  
CC tumouricidal immunocyte population, and administering the tumouricidally  
CC activated immunocytes to the host), producing (M3) a tumouricidal APC  
CC population ex vivo in a mammal (by allowing a tumour associated lipid to  
CC contact APCs, in which receptors for the tumour associated lipids are  
CC inactivated or deleted to produce a tumouricidally activated population,  
CC and administering APCs to the host), producing a tumouricidal T cell  
CC population ex vivo in a mammal) by allowing a tumour associated lipids to  
CC contact T cells, in which adaptor proteins, which inhibit T cell  
CC activation by tumour associated antigens, are deleted or functionally  
CC deactivated to produce a tumouricidal population of T cells, and  
CC administering the tumouricidally activated T cells to the host, or  
CC allowing a superantigen-lipid raft to contact T cells ex vivo, and  
CC administering the tumouricidally activated T cells to the host), treating  
CC (M5) cancer in a mammal (by administering a lipid binding molecule which  
CC binds immunosuppressive tumour associated lipids in vivo), producing (M6)  
CC a tumouricidal T cell population in vivo in a mammal (by allowing a  
CC tumour associated antigen to contact immunocytes in which adaptor  
CC proteins which inhibit T cell activation by tumour associated antigens  
CC are deleted or functionally deactivated) and producing (M7) a  
CC tumouricidal T cell population ex vivo in a mammal comprising allowing a  
CC superantigen-lipid raft conjugate to contact immunocytes in vivo. The  
CC receptors, methods and compositions are useful for treating cancers and  
CC tumours. Bacterial superantigens are co-administered or administered as  
CC fusion constructs with anti-tumour proteins or motifs. The present  
CC sequence represents an anti-tumour protein which is co-administered with

CC or incorporated into a fusion construct with a superantigen. Note: The  
CC sequence data for this patent did not form part of the printed  
CC specification, but was obtained in electronic format from the US patent  
CC office website at "seqdata.uspto.gov/sequence.html?DocID=20020177551"  
XX  
XX  
SQ Sequence 1183 AA;  
  
Query Match 93.4%; Score 1048; DB 6; Length 1183;  
Best Local Similarity 100.0%; Pred. No. 1.4e-78;  
Matches 200; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
  
QY 12 SDDKVATITSGNKSTNVTYHKSAGTSSVFYKTKGMDLPEDTTHVRFLNINNEKSYVSK 71  
Db 144 SDDKVATITSGNKSTNVTYHKSAGTSSVFYKTKGMDLPEDTTHVRFLNINNEKSYVSK 203  
QY 72 DITIKQIQGGQQLDLSTLNVGTHTSNYSYQSASITDFEKAPPGSKITVDNTKNTIDV 131  
Db 204 DITIKQIQGGQQLDLSTLNVGTHTSNYSYQSASITDFEKAPPGSKITVDNTKNTIDV 263  
QY 132 TIPOGYSYNSFSINYKTKITNEQOKEFVNNSQAQYOEHGKKEEVNGKSFNHTVHNINANA 191  
Db 264 TIPOGYSYNSFSINYKTKITNEQOKEFVNNSQAQYOEHGKKEEVNGKSFNHTVHNINANA 323  
QY 192 GIEGTVKGLKVLKQDKDTK 211  
Db 324 GIEGTVKGLKVLKQDKDTK 343  
  
RESULT 4  
AAR22675  
ID AAR22675 standard; protein; 1185 AA.  
XX  
XX AAR22675;  
XX  
DT 25-MAR-2003 (revised)  
DT 04-DEC-1992 (first entry)  
XX  
DE Collagen binding protein.  
XX  
KW CBP; collagen binding protein; mastitis; arthritis.  
XX  
OS Staphylococcus aureus.  
XX  
FH Key Location/Qualifiers  
FT Peptide 1..29 /label= sig\_peptide  
FT Region 30..1185 /note= "claim 11; page 29-30"  
FT Region 30..534 /label= A  
FT Region 535..721 /note= "see CC"  
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FT Region 10780..10799 /note= "see CC"  
FT Region 10800..10819 /note= "see CC"  
FT Region 10820..10



CC infections, and in production facilities or laboratories to isolate  
CC additional quantities of collagen-binding proteins. It is also useful for  
CC treating medical instruments in order to reduce or eliminate the  
CC possibility of them becoming infected or further spreading the infection.  
CC The present sequence is *S. aureus* CNA19 protein  
XX  
SQ Sequence 168 AA;  
  
Query Match 78.8%; Score 884; DB 4; Length 168;  
Best Local Similarity 99.4%; Pred. No. 5.3e-66;  
Matches 167; Conservative 0; Mismatches 1; Indels 0; Gaps 0;  
  
Qy 19 ITSQKSTNTVHKSEAGTSVFYKTKGDMLPEDTHVRWFLNINNEKSYVKDITIKDQ 78  
Db 1 ITSQKSTNTVHKSEAGTSVFYKTKGDMLPEDTHVRWFLNINNEKSYVKDITIKDQ 60  
  
Qy 79 IQGQQQLDLSTLNINVTGTHSNYSGQSAITDFKAPPGSKIIVDNTKNTIDVTIPQY 138  
Db 61 IQGQQQLDLSTLNINVTGTHSNYSGQSAITDFKAPPGSKIIVDNTKNTIDVTIPQY 120  
  
Qy 139 SYNSFSINYKTKITNEQKQEFVNNSQAWYQEHGKEEVNGKSFNHTVHN 186  
Db 121 SYNSFSINYKTKITNEQKQEFVNNSQAWYQEHGKEEVNGKSFNHTVHN 168  
  
RESULT 6  
AAW31552  
ID AAW31552 standard; protein; 159 AA.  
XX  
AC AAW31552;  
XX  
DT 27-AUG-2003 (revised)  
DT 25-MAR-2003 (revised)  
DT 21-MAY-1998 (first entry)  
XX  
DE Collagen binding protein M17 epitope.  
XX  
KW Collagen binding protein; cna gene; sepsis; infection;  
KW microbial surface component recognising adhesive matrix molecule; MSCRAMM;  
KW adhesin; vaccine; immunisation; diagnosis; therapy; epitope M17.  
XX  
OS Staphylococcus aureus.  
XX  
FH Key Location/Qualifiers  
FT Peptide 1..12  
FT /note= "vector pQE30-derived peptide"  
FT Protein 13..159  
FT /note= "epitope M17"  
XX  
FN WO9743314-A2.  
XX  
XX 20-NOV-1997.  
XX  
XX 14-MAY-1997; 97WO-US008210.  
XX  
XX 16-MAY-1996; 96US-0017678P.  
XX  
XX (TEXA ) UNIV TEXAS A & M SYSTEM.  
XX (UABR-) UAB RES FOUND.  
XX  
XX Hoocek M, Patti JM, House-Pompeo K, Sthanam N, Symersky J;  
XX  
XX WPI; 1998-008801/01.  
XX N-PSDB; AAT93436.  
XX  
XX Antibody that interacts with collagen binding domain of Staphylococcal  
XX cna gene product - useful to prevent bacterial sepsis in animal infected  
XX with Staphylococcus aureus.  
XX  
XX Claim 31; Page 114; 143pp; English.  
XX  
XX This protein comprises Staphylococcus aureus collagen binding protein  
CC (CBP) epitope M17, i.e. amino acids 151-297 of full-length CBP, plus a

CC vector-derived N-terminal peptide. Claimed 441, 849 and 1500 bp nucleic  
CC acid sequences (see AAT93436-38) respectively encode CBP epitopes M17,  
CC M31 and M55 (see AAW31552-54) that confer protection against *S. aureus*  
CC infection. These nucleic acid sequences can be used in the recombinant  
CC production of the CBP epitopes. The CBP protein and antigenic epitopes  
CC are contemplated for use in the treatment of pathological infections,  
CC especially to prevent bacterial adhesion to collagen. The claimed nucleic  
CC acids as well as claimed anti-CBP antibodies will also be of use in  
CC screening, diagnostic and therapeutic applications including active and  
CC passive immunisation and methods for the prevention of bacterial  
CC colonisation in an animal such as a human. The CBP epitopes are also  
CC contemplated for use in the preparation of vaccines and as carrier  
CC proteins in vaccine formulations, as well as in the formulation of  
CC compositions for the prevention of *S. aureus* infection. (Updated on 25-  
CC MAR-2003 to correct PI field.) (Updated on 27-AUG-2003 to correct OS  
CC field.)  
XX  
SQ Sequence 159 AA;  
  
Query Match 74.0%; Score 830; DB 2; Length 159;  
Best Local Similarity 96.4%; Pred. No. 1.6e-61;  
Matches 159; Conservative 0; Mismatches 0; Indels 6; Gaps 1;  
  
Qy 1 MRGSHHHHGGSDDKVATITSGNKSNTVTYHKSEAGTSVFYKTKGDMLPEDTHVRWFL 60  
Db 1 MRGSHHHHGGSDDKVATITSGNKSNTVTYHKSEAGTSVFYKTKGDMLPEDTHVRWFL 54  
  
Qy 61 NINNEKSYVSKDITIKDQIQGGQQLDLSTLNINVTGTHSNYSGQSAITDFKAPPGSKI 120  
Db 55 NINNEKSYVSKDITIKDQIQGGQQLDLSTLNINVTGTHSNYSGQSAITDFKAPPGSKI 114  
  
Qy 121 TVDNTKNTIDVTIPQGYGYSNFSINFKTKITNEQKQEFVNNSQA 165  
Db 115 TVDNTKNTIDVTIPQGYGYSNFSINFKTKITNEQKQEFVNNSQA 159  
  
RESULT 7  
ADC97372  
ID ADC97372 standard; protein; 731 AA.  
XX  
AC ADC97372;  
XX  
DT 01-JAN-2004 (first entry)  
XX  
DE E. faecium protein sequence SEQ ID 6999.  
XX  
KW Vaccine; urinary tract infection; bacteraemia; endocarditis; wound;  
KW abdominal-pelvic infection.  
XX  
OS Enterococcus faecium.  
XX  
XX US6583275-B1.  
XX  
XX 24-JUN-2003.  
XX  
XX 30-JUN-1998; 98US-00107532.  
XX  
XX 02-JUL-1997; 97US-0051571P.  
XX 14-MAY-1998; 98US-0085598P.  
XX  
XX (GENO-) GENOME THERAPEUTICS CORP.  
XX  
XX Doucette-Stamm LA, Bush D;  
XX  
XX WPI; 2003-799836/75.  
XX N-PSDB; ADC93718.  
XX  
XX New isolated nucleic acid derived from Enterococcus faecium encoding an  
XX Enterococcus faecium polypeptide useful for detection, prevention and  
XX treatment of a pathological condition resulting from a bacterial  
XX infection.  
XX  
XX Example 1; SEQ ID NO 6999; 243pp; English.

XX The invention relates to an isolated nucleic acid derived from  
 CC Enterococcus faecium encoding an Enterococcus faecium polypeptide having  
 CC one of 10 fully defined sequences given in the (or comprising 40  
 CC sequential nucleotides chosen from any of the nucleic acids, its  
 CC complement or sequences hybridising to it). Also included are a  
 CC recombinant vector comprising the nucleic acid operably linked to  
 CC transcription regulatory element, a cell comprising the vector and a  
 CC single-stranded probe comprising the nucleic acid. The nucleic acids are  
 CC chosen from 3654 disclosed sequences encoding 3654 disclosed proteins.  
 CC The nucleic acids is useful for diagnosing pathological conditions  
 CC resulting from E. faecium bacterial infection (e.g. urinary tract  
 CC infection, bacteraemia, endocarditis, wounds and abdominal-pelvic  
 CC infection) and for screening drugs such as agonists and antagonists. The  
 CC nucleic acid is useful for recombinant production of Candida albicans -  
 CC derived peptides or antisense polypeptides. Pharmaceutical compositions  
 CC and vaccines containing the nucleic acid are useful for preventing or  
 CC treating Enterococcus faecium infections. The present sequence represents  
 CC one if the disclosed E. faecium proteins.  
 XX  
 SQ Sequence 731 AA;

Query Match 43.8%; Score 491; DB 7; Length 731;  
 Best Local Similarity 48.5%; Pred. No. 2.6e-32; Indels 2; Gaps 2;  
 Matches 95; Conservative 35; Mismatches 64;  
 QY 18 TITSGNKSTNVTVHKSEAG-TSSVFYKTDGMLPDTTHVRWFLNINNEKSYVSKDIT 76  
 DB 162 TVTSGDKATVNTVKPASGSSSVFYKTDGMLPDTKHIRWFLNINNGTYVEQPKIS 221  
 QY 77 DQIQGGQQLDLSTLIN-VTGTHSNYYSGSAITDFEKAFFPGSKITVDNTKNTIDVTIPQ 135  
 DB 222 DEIQQGQQLDPSTFEINQIHLGEQKYRBEIGIQQLDFPSATFNFSVDNIEITIPK 281  
 QY 136 GYGSYNSFSINYKTKITNEQKFEVNSQAWYQEHGKEEVNGKSFNHTVHNINANAGIEG 195  
 DB 282 NFVNLKIMVSYKTIENPQINFNHSEAWFEKFNKPAVDGSEFNHTVKNISAGSGVNG 341  
 QY 196 TVRGELKVLKQDKDTK 211  
 DB 342 TVRGELKIFKVINDE 357

RESULT 8  
 AAB37667  
 ID AAB37667 standard; protein; 458 AA.

XX AAB37667;  
 AC AAB37667;  
 XX 28-FEB-2001 (first entry)  
 DT Collagen binding Ace protein.  
 DE Collagen binding Ace protein.  
 XX Collagen binding Ace protein; antibacterial; immunostimulant; vaccine;  
 KW extracellular matrix-binding protein; lamin binding; bacterial infection.  
 XX Enterococcus faecalis.  
 OS  
 XX WO2000068242-A1.  
 PN  
 XX 16-NOV-2000.  
 PD  
 XX 10-MAY-2000; 2000WO-US012590.  
 PF  
 XX 10-MAY-1999; 99US-0133334P.  
 PR  
 XX (TEXA ) UNIV TEXAS A & M SYSTEM.  
 PA (TEXA ) UNIV TEXAS MEDICAL SCHOOL.  
 XX Rich RL, Kriekemeyer B, Owens RT, Hook M, Murray BE;  
 PI Nallapareddy SR, Qin X, Weinstock GM, Singh KV, Duh R;  
 XX WPI; 2000-687639/67.

DR N-PSDB; AAC83828.  
 XX New collagen-binding protein from Enterococcus, useful e.g. in protective  
 PT vaccines, for diagnosis and treatment of Enterococcal infections and for  
 PT screening for compounds that inhibit collagen binding by enterococci.  
 XX  
 PS Claim 26; Page 119-122; 148pp; English.  
 XX  
 CC The present sequence is collagen binding Ace protein from Enterococcus  
 CC faecalis, where Ace is adhesion of collagen from enterococcal bacteria.  
 CC This protein is an extracellular matrix-binding protein, which can bind  
 CC with collagens such as collagen type I and type IV and with lamins. Ace  
 CC protein can be used to inhibit attachment of enterococci to collagen and  
 CC to treat or prevent enterococcal infections, specifically as a vaccine  
 XX  
 SQ Sequence 458 AA;

Query Match 15.9%; Score 178; DB 3; Length 458;  
 Best Local Similarity 28.3%; Pred. No. 2.1e-06; Indels 16; Gaps 6;  
 Matches 54; Conservative 33; Mismatches 88;  
 QY 18 TITSGNKSTNVTVHKSEAG-TSSVFYKTDGMLPDTTHVRWFLNINNEKSYVSKDIT 74  
 DB 121 TATATQRLTIEGVNTETQIERDYFFYKVGDLAGE-SNQVRWFLNVLNKSVDIEDIS 179  
 QY 75 IKDQIQGGQQLDLSTLIN-VTGTHSNYYSGSAITDFEKAFFPGSKITVDNTKNTIDVTIP 134  
 DB 180 IADRQSGQQLNKESFTFDIVNDKETKY---ISLAEFEQGGYKIDFV--TDNDFNLRFY 234  
 QY 135 QYGSYNSFSINYKTKIT-NEQKFEVNSQAWYQEHGKEEVNGKSFNHTVHNINANAG 192  
 DB 235 RKAREFTSFIVRYTSTITEAGHQATFENSYDINQLNNQDATNEKTSQV-----KNVF 289  
 QY 193 IEGTVKGLKV 203  
 DB 290 VEGEASGNQNV 300

RESULT 9  
 ADD90648  
 ID ADD90648 standard; protein; 458 AA.

XX ADD90648;  
 XX 29-JAN-2004 (first entry)  
 DT Enterococcus faecalis collagen binding ACE protein SEQ ID NO:1.  
 DE Enterococcus faecalis collagen binding ACE protein SEQ ID NO:1.  
 XX cross-reactive monoclonal antibody; ACE; Enterococcus faecalis;  
 KW antibacterial; veterinary; antibody therapy; vaccine; infection;  
 KW collagen binding ACE.  
 XX Enterococcus faecalis.  
 OS  
 XX WO2003072607-A1.  
 PN  
 XX 04-SEP-2003.  
 PD  
 XX 21-FEB-2003; 2003WO-US005040.  
 PF  
 XX 21-FEB-2002; 2002US-0357832P.  
 PR  
 XX 05-MAR-2002; 2002US-0361347P.  
 XX (UYPA-) UNIV PAVIA.  
 PA  
 XX Bates SE, Speziale P, Visai L, Giampiero P;  
 PI WPI; 2003-679944/64.  
 XX N-PSDB; ADD90649.  
 DR  
 XX New cross-reactive monoclonal antibody that binds to a peptide such as  
 PT the A domain of the ACE protein from Enterococcus faecalis, or CAN19 from  
 PT Staphylococcus aureus, useful for treating or preventing infections

PT caused by the organisms.

PS Claim 8; SEQ ID NO 1; 49pp; English.

XX  
XX The present invention describes a cross-reactive monoclonal antibody (I)  
CC that binds to a peptide selected from the A domain of the ACE protein  
CC from Enterococcus faecalis, ACE 19 from E. faecalis and CAN19 from  
CC Staphylococcus aureus. Also described: (1) an isolated antisera  
CC containing binding (I); (2) a diagnostic kit comprising (I) and means for  
CC detecting binding by (I); (3) a pharmaceutical composition for treating  
CC or preventing a bacterial infection comprising (I), and a vehicle,  
CC carrier or excipient; (4) a method for treating or preventing an  
CC infection of enterococcal, streptococcal, or staphylococcal infection by  
CC administering (I) to a human or animal patient; and (5) a method for  
CC inducing an immunological response by administering to human or animal  
CC an immunogenic amount of an isolated protein from E. faecalis selected  
CC from ACE40 and ACE19 protein. (I) has antibacterial and veterinary  
CC activities, and can be used in antibody therapy, and in vaccines. The  
CC antibody and pharmaceutical composition are useful for treating or  
CC preventing infection from staphylococcal, streptococcal and enterococcal  
CC bacteria in a human or animal. The present sequence represents the  
CC Enterococcus faecalis collagen binding ACE protein, which is used in the  
CC exemplification of the present invention.

XX SQ Sequence 458 AA;

Query Match 15.9%; Score 178; DB 7; Length 458;

Best Local Similarity 28.3%; Pred. No. 2.1e-06;

Matches 54; Conservative 33; Mismatches 88; Indels 16; Gaps 6;

QY 18 TITSGNKSNTVTHKSAG--TSSVFFYKTDMLPETHVRWFLNINNEKSYVSKDIT 74

DB 121 TATATQRLTIEGVTNTGTQIERDYPFFYKVGDLAGE-SNQVRWFLNINNEKSDVTEDIS 179

QY 75 IKQIQGGQQLDLSTLNINVTGTHSNYSQSAITDFEKAPGSKITVDNKTIDVTIP 134

DB 180 IADRQGGQQLNKESTFTDINDKETYK---ISLAEEFQGGYKIDFV--TDNDFNLRFY 234

QY 135 QGYGSYNSFSINYTKKIT--NEOQKEFVNSQAWYQHGKKEEVNKGSKFNHTVHNINANAG 192

DB 235 RDKARFTSFIVRYTSTITEAGHQHATFENSVDYINQLNNQDATNEKTSQV-----KNVF 289

QY 193 IEGTVKGELKV 203

DB 290 VEGEASGNQNV 300

RESULT 10

AAB37668

ID AAB37668 standard; protein; 146 AA.

AC AAB37668;

XX 28-FEB-2001 (first entry)

DE Collagen binding Ace protein collagen binding domain.

DE Collagen binding Ace protein; antibacterial; immunostimulant; vaccine;

KW extracellular matrix-binding protein; lamin binding; bacterial infection.

KW Enterococcus faecalis.

OS WO200068242-A1.

XX 16-NOV-2000.

XX 10-MAY-2000; 2000WO-US012590.

XX 10-MAY-1999; 99US-0133334P.

XX (TEXA ) UNIV TEXAS A & M SYSTEM.

PA (TEXA ) UNIV TEXAS MEDICAL SCHOOL.

XX

PI Rich RL, Kriekemeyer B, Owens RT, Hook M, Murray BE;  
PI Nallapareddy SR, Qin X, Weinstock GM, Singh KV, Duh R;  
XX WPI; 2000-687639/67.

XX New collagen-binding protein from Enterococcus, useful e.g. in protective  
PT vaccines, for diagnosis and treatment of Enterococcal infections and for  
PT screening for compounds that inhibit collagen binding by enterococci.

XX Disclosure; Fig 1; 148pp; English.

XX The present invention relates to collagen binding Ace protein from  
CC Enterococcus faecalis, where Ace is adhesion of collagen from  
CC enterococcal bacteria (see AAC83828 and AAB37667). Ace protein is an  
CC extracellular matrix-binding protein, which can bind with collagens such  
CC as collagen type I and type IV and with lamins. Ace protein can be used  
CC to inhibit attachment of enterococci to collagen and to treat or prevent  
CC enterococcal infections, specifically as a vaccine. The present sequence  
CC is the collagen binding domain of Ace protein

XX SQ Sequence 146 AA;

Query Match 14.8%; Score 165.5; DB 3; Length 146;

Best Local Similarity 30.2%; Pred. No. 5e-06;

Matches 45; Conservative 30; Mismatches 65; Indels 9; Gaps 5;

QY 41 FYYKTGDMLPEDTHVRWFLNINNEKSYVSKDITIKDQIQGGQQLDLSTLNINVTGTHSN 100

DB 5 FFFKVGDLAGE-SNQVRWFLNINNEKSDVTEDISIADRQGGQQLNKESTFTDINDKET 63

QY 101 YYGQSAITDFEKAPGSKITVDNKTIDVTIPQGYGSYNSFSINYTKKIT--NEOQKE 158

DB 64 KY---ISLAEEFQGGYKIDFV--TDNDFNLRFYRDKARFTSFIVRYTSTITEAGHQHAT 118

QY 159 FVNSQAWYQHGKKEEVNKGSKFNHTVHNI 187

DB 119 FENSVDYINQLNNQDATNEKTSQ-VKNV 146

RESULT 11

AAE11856

ID AAE11856 standard; protein; 168 AA.

AC AAE11856;

XX 18-DEC-2001 (first entry)

DE Enterococcus faecalis ACE19 protein.

DE Collagen-binding region; CNA19; Staphylococcus aureus infection;

KW Staphylococcus epidermidis infection; vaccine; CNA protein;

KW cross-reactive antibody; collagen-adhesin; ACE.

OS Enterococcus faecalis.

XX WO200170267-A1.

XX 27-SEP-2001.

XX 19-MAR-2001; 2001WO-US008554.

XX 17-MAR-2000; 2000US-0189968P.

XX 25-APR-2000; 2000US-0199370P.

XX 15-AUG-2000; 2000US-0225402P.

XX (INHI-) INHIBITEX INC.

PA (TEXA ) UNIV TEXAS A & M SYSTEM.

PA (UYPA-) UNIV PAVIA.

XX Hook M, Xu Y, Speciale P, Visai L, Casolini F, Patti J, Patel P;

PI Domanski P;

XX WPI; 2001-607512/69.





PA (INH1-) INHIBITEX INC.  
 XX (TEXA ) UNIV TEXAS A & M SYSTEM.  
 PI Patti JM, Hutchins JT, Hall A, Domanski P, Patel P, Hook M;  
 PI Robbins J, Vernachio J, Bowden MG;  
 XX  
 XX  
 DR WPI; 2003-722324/68.  
 DR N-PSDB; ACF80625.  
 XX  
 XX New antibody recognizing a Staphylococcus epidermidis protein comprising  
 PT SdrG N1N2N3, SdrG N2N3 or SdrGTR2 useful for preparing a composition for  
 PT treating or preventing a coagulase-negative Staphylococcal infection.  
 XX  
 XX Claim 27; Page 25; 78pp; English.  
 XX  
 XX The present sequence comprises the protein sequence of the N2N3 region  
 CC (amino acids 273-597) of the SdrG surface protein of coagulase-negative  
 CC Staphylococcus epidermidis. A claimed antibody recognises a protein  
 CC selected from SdrG N1N2N3, N2N3 and TR2. The antibody may be a monoclonal  
 CC antibody, including a chimeric, murine, humanized, human or single chain  
 CC monoclonal antibody, which prevents a coagulase-negative staphylococcal  
 CC infection in a human or animal by inhibiting binding of staphylococcus  
 CC bacteria to fibrinogen. Such antibodies can be used to treat or prevent  
 CC staphylococcal infections including nosocomial coagulase-negative  
 CC staphylococcal infections in low birth weight infants. A claimed vaccine  
 CC comprises the isolated SdrG N1N2N3, N2N3 or TR2 protein  
 XX  
 XX Sequence 343 AA;  
 Query Match 11.7%; Score 131.5; DB 7; Length 343;  
 Best Local Similarity 25.3%; Pred. No. 0.011; Indels 85; Gaps 16;  
 Matches 66; Conservative 32; Mismatches 78; Indels 85; Gaps 16;  
 QY 1 MRGSHHHHHHGS-----DDKVATITSGNKSTN--VTVHKSEAGTSSV 40  
 DB 1 MRGSHHHHHHGSVPRGSEQSNVNLKVTQD--SITEGYDDSDGIKAHDAE---NL 54  
 QY 41 PY---YKTDMLPBDTHVRWFLNINNEKSVSKDIT-----IKDQIOGGQQLDLST- 89  
 DB 55 IYDVTFFVDKVKSGDT-----MTVNIDKNTVPSDLTDSFAIPKIKD--NSGEIATGTY 107  
 QY 90 --LNINVTGTHSNY---YSGQSAITDFEAPGSKITVDNTKNTIDVTIPQGYGSYNSFS 144  
 DB 108 DNTWKQITYTFTDVKYENIKAHKLTSYDKSKVPNNNTK--LDVEYKLTALSSVN--- 162  
 QY 145 INYKTKITNEQKEFVN---NSQAWYQEHGKEEVNGKSFNHTVH-----TNIDTKHTVEQTYINPLYSAKETN 185  
 DB 163 ---KT-ITVEYQKPNENRTANLQSWF-----TNIDTKHTVEQTYINPLYSAKETN 211  
 QY 186 -NINAGIEGTVKGLKVLK 205  
 DB 212 VNISGNGDEGSTIIDSTIIK 232  
 RESULT 14  
 AAW31555  
 ID AAW31555 standard; protein; 345 AA.  
 XX  
 XX AAW31555;  
 XX  
 XX 27-AUG-2003 (revised)  
 DT 25-MAR-2003 (revised)  
 DT 21-MAY-1998 (first entry)  
 XX  
 XX Fibronectin-binding MSCRAMM derivative pCF33.  
 XX  
 XX Fibronectin; pCF33; collagen binding protein; sepsis; infection;  
 KW microbial surface component recognising adhesive matrix molecule; MSCRAMM;  
 KW adhesin; vaccine; immunisation; diagnosis; therapy; mastitis.  
 XX  
 XX Staphylococcus aureus.  
 OS  
 XX  
 XX Key Location/Qualifiers  
 FH

Peptide 1. .12  
 /note= "vector pOE30-derived peptide"  
 FT  
 XX WO97433314-A2.  
 XX  
 XX 20-NOV-1997.  
 PD  
 XX  
 XX 14-MAY-1997; 97WO-US008210.  
 PF  
 XX  
 XX 16-MAY-1996; 96US-0017678P.  
 PR  
 XX  
 XX (TEXA ) UNIV TEXAS A & M SYSTEM.  
 PA  
 PA (UABR-) UAB RES FOUND.  
 XX  
 XX Hoeok M, Patti JM, House-Pompeo K, Sthanam N, Symersky J;  
 PI  
 PI WPI; 1998-008801/01.  
 DR  
 XX  
 XX Antibody that interacts with collagen binding domain of Staphylococcal  
 PT cna gene product - useful to prevent bacterial sepsis in animal infected  
 PT with Staphylococcus aureus.  
 XX  
 XX Disclosure; Page 91; 143pp; English.  
 PS  
 XX  
 XX This protein comprises Staphylococcus aureus fibronectin-binding  
 CC microbial surface component recognising adhesive matrix molecule (MSCRAMM)  
 CC derivative pCF33, plus a vector-derived N-terminal peptide. The invention  
 CC relates to claimed nucleic acid sequences (see AAT93436-38) encoding S.  
 CC aureus collagen binding protein (CBP) epitopes M17, M31 and M55 (see  
 CC AAW31552-54) that confer protection against S. aureus infection. CBP  
 CC protein and antigenic epitopes are contemplated for use in the treatment  
 CC of pathological infections, especially to prevent bacterial adhesion to  
 CC collagen. The epitopes are also contemplated for use in the preparation  
 CC of vaccines and as carrier proteins in vaccine formulations, as well as  
 CC in the formulation of compositions for the prevention of S. aureus  
 CC infection. pCF33 and PQD (see AAW31556) were used to raise anti-MSCRAMM  
 CC polyclonal antibodies used in passive immunisation against bovine  
 CC mastitis. (Updated on 25-MAR-2003 to correct PI field.) (Updated on 27-  
 CC AUG-2003 to correct OS field.)  
 XX  
 XX Sequence 345 AA;  
 Query Match 11.2%; Score 125.5; DB 2; Length 345;  
 Best Local Similarity 30.1%; Pred. No. 0.035;  
 Matches 56; Conservative 20; Mismatches 73; Indels 37; Gaps 10;  
 QY 1 MRGSHHHHHHGS---DDKVATITSGNKSTN-----TVHKSEAGTSSVYKGTGDM 48  
 DB 1 MRGSHHHHHHGSVMADAPAGTDTITNQLTNVTGDSGTTVPHQAGYKLN---GFS 57  
 QY 49 LPEDTTHVRWF-----LNINNEKSVS-KDITIKDQIOGGQQLDLSTLNINVTGTHSN 100  
 DB 58 VPNSAVKGDFTFKITVPKELNLTGVTSPAKVPPIMAGDQVLANGVIDSDG---NVIYTFD 114  
 QY 101 YSGQSAITDFEKA---FPGSKITVDNTKNTIDVTIPQGYGSYNSFINKTKITNEQOK 157  
 DB 115 YVNTK-----DDVKATLTMP-AYIDPENVKKTGNVTATGICGTTA---NKTVLVDYKEYG 166  
 QY 158 EFVNNNS 163  
 DB 167 KFYNLS 172  
 RESULT 15  
 ABB53995  
 ID ABB53995 standard; protein; 336 AA.  
 XX  
 XX ABB53995;  
 XX  
 XX 29-AUG-2003 (revised)  
 DT 16-MAY-2002 (first entry)  
 XX  
 XX Lactococcus lactis protein icaC.

XX Biosynthesis; biodegradation; lactic bacterium; yogurt; cheese.  
XX Lactococcus lactis; IL1403.  
XX FR2807446-A1.  
XX 12-OCT-2001.  
XX 11-APR-2000; 2000FR-00004630.  
XX 11-APR-2000; 2000FR-00004630.  
XX (INRG ) INRA INST NAT RECH AGRONOMIQUE.  
XX Bolotine A, Sorokine A, Renault P, Ehrlich SD;  
XX WPI; 2002-043418/06.  
XX New nucleotide sequence useful in the identification or Lactococcus  
XX lactis and related species.  
XX Claim 6; SEQ ID NO 697; 2504pp; French.  
XX The present invention is related to a Lactococcus lactis nucleotide  
XX sequence (ABA90521) and related proteins (ABBS3300-ABBS5621). The nucleic  
XX acid sequence is useful in the detection and/or amplification of nucleic  
XX acid sequence, particularly to identify Lactococcus lactis or related  
XX species. The proteins of the invention are useful for the biosynthesis or  
XX biodegradation of a composition of interest. The invention helps research  
XX in lactic bacteria, particularly useful in the production of yogurt and  
XX cheese. Note: The sequence data for this patent is based on equivalent  
XX patent WO200177334 (published 18-OCT-2001) which is available in  
XX electronic format directly from WIPO at  
XX fip.wipo.int/pub/published\_pdt\_sequences. (Updated on 29-AUG-2003 to  
XX standardise OS field)  
XX SQ Sequence 336 AA;

Query Match 10.6%; Score 119; DB 5; Length 336;  
Best Local Similarity 25.2%; Pred. No. 0.12; Indels 58; Gaps 11;  
Matches 52; Conservative 28; Mismatches 68;  
Qy 24 KSTNVTYHKEAGTSSV-----FYKTGDMLPEDTTHVRWFLNNKESYVSK 71  
Db 71 EDTRVVDWKDGGTTEVNI GPATGPKDEVLYKMGWVDENDSTLIHWQVRINYAKENIQK 130  
Qy 72 DITIKDOIQQOQL---DLSTLINVTGTHSNY----YSGQSAITDFEKAPPGSKITVDN 124  
Db 131 AI-YTDIIGGNQNLVSGSISVANVTYSSDGENYVDSYFQASI--LENGVNGFTVNLGD 187  
Qy 125 TKNTIDVTIPQYGSYNSFSINVKITN---EQCKEFVNNSQAWYQEHGKEEVNGKSFN 181  
Db 188 ISNTI-----IIDYSTRATDGLSQ-----YENRG--ELTGENIE 221  
Qy 182 -----HTVHN-INANAGIEGTVKE 200  
Db 222 KQVVEVHTPNNGNGNASKLSISGE 247

Search completed: July 27, 2004, 10:08:19  
Job time : 57 secs

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OM protein - protein search, using sw model

Run on: July 27, 2004, 10:07:21 ; Search time 19 Seconds  
(without alignments)  
573.320 Million cell updates/sec

Title: US-09-813-820-4  
Perfect score: 1122  
Sequence: 1 MRGSHHHHSGDDKVATIT.....GIEGTGKELKVLKQDKTK 211

Scoring table: BLOSUM62  
Gapop 10.0 , Gapext 0.5

Searched: 389414 seqs, 51625971 residues

Total number of hits satisfying chosen parameters: 389414

Minimum DB seq length: 0  
Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%  
Maximum Match 100%  
Listing first 45 summaries

Database : Issued\_Patents\_AA:\*  
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2: /cgn2\_6/prodata/2/iaa/5B\_COMB.pep.\*  
3: /cgn2\_6/prodata/2/iaa/6A\_COMB.pep.\*  
4: /cgn2\_6/prodata/2/iaa/6B\_COMB.pep.\*  
5: /cgn2\_6/prodata/2/iaa/PCTUS\_COMB.pep.\*  
6: /cgn2\_6/prodata/2/iaa/backfiles1.pep.\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	ID	Description
1	1132	100.0	211	3	US-08-856-253-4
2	1054.5	94.0	512	3	US-08-856-253-6
3	1048	93.4	1183	2	US-08-447-031A-2
4	830	74.0	159	3	US-08-856-253-2
5	491	43.8	731	4	US-09-107-532A-6999
6	178	15.9	540	4	US-09-134-000C-3745
7	125.5	11.2	345	3	US-08-856-253-7
8	114.5	10.2	688	4	US-09-071-035-464
9	114.5	10.2	2032	4	US-09-071-035-458
10	114.5	10.2	2032	4	US-09-071-035-462
11	114.5	10.2	2032	4	US-09-071-035-466
12	114.5	10.2	2054	4	US-09-134-000C-6612
13	113.5	10.1	184	3	US-09-023-082A-30
14	113.5	10.1	184	4	US-09-248-998-30
15	110.5	9.8	1338	2	US-08-728-470-9
16	110.5	9.8	1338	3	US-08-719-641-9
17	110.5	9.8	1599	2	US-08-617-697-9
18	110	9.8	2314	4	US-09-268-347-49
19	104.5	9.3	1529	3	US-08-728-470-10
20	104.5	9.3	1529	3	US-08-719-641-10
21	104.5	9.3	1600	2	US-08-617-697-10
22	103.5	9.2	139	3	US-08-856-253-8
23	103.5	9.2	523	4	US-09-543-681A-5151
24	100	8.9	1095	4	US-09-206-942-69
25	100	8.9	1536	1	US-08-038-682-2
26	100	8.9	1536	1	US-08-302-832-2
27	100	8.9	1536	2	US-08-530-198-2

28	100	8.9	1536	2	US-08-469-880-2	Sequence 2, Appli
29	100	8.9	1536	2	US-08-728-470-2	Sequence 2, Appli
30	100	8.9	1536	2	US-08-617-697-2	Sequence 2, Appli
31	100	8.9	1536	3	US-08-719-641-2	Sequence 2, Appli
32	100	8.9	1536	4	US-09-206-942-67	Sequence 67, Appl
33	99.5	8.9	589	3	US-09-377-155-19	Sequence 19, Appl
34	99.5	8.9	589	4	US-09-669-974-19	Sequence 19, Appl
35	99.5	8.9	589	4	US-09-797-862-19	Sequence 19, Appl
36	98.5	8.8	351	2	US-08-701-191A-3	Sequence 3, Appli
37	98.5	8.8	351	4	US-09-664-526-3	Sequence 3, Appli
38	96.5	8.6	1742	4	US-09-386-962C-4	Sequence 4, Appli
39	95.5	8.5	1436	4	US-10-080-505-13	Sequence 13, Appl
40	95	8.5	666	2	US-08-737-716-14	Sequence 14, Appl
41	94.5	8.4	514	4	US-09-107-532A-4411	Sequence 4411, Ap
42	94	8.4	287	4	US-09-543-681A-5491	Sequence 5491, Ap
43	94	8.4	821	4	US-09-556-877-195	Sequence 195, App
44	94	8.4	821	4	US-09-620-412C-195	Sequence 195, App
45	94	8.4	821	4	US-09-598-419-195	Sequence 195, App

ALIGNMENTS

RESULT 1  
US-08-856-253-4  
; Sequence 4, Application US/08856253  
; Patent No. 6288214  
; GENERAL INFORMATION:  
; APPLICANT: Hook, Magnus  
; APPLICANT: Patti, Joseph M.  
; APPLICANT: House-Pompeo, Karen  
; APPLICANT: Sthanam, Narayana  
; APPLICANT: Symersky, Jindrich  
; TITLE OF INVENTION: COLLAGEN BINDING PROTEIN COMPOSITIONS  
; TITLE OF INVENTION: AND METHODS OF USE  
; NUMBER OF SEQUENCES: 8  
; CORRESPONDENCE ADDRESS:  
; ADDRESSEE: Arnold, White & Durkee  
; STREET: P.O. Box 4433  
; CITY: Houston  
; STATE: Texas  
; COUNTRY: U.S.  
; ZIP: 77210  
; COMPUTER READABLE FORM:  
; MEDIUM TYPE: Floppy disk  
; COMPUTER: IBM PC compatible  
; OPERATING SYSTEM: PC-DOS/MS-DOS  
; SOFTWARE: PatentIn Release #1.0, Version #1.30  
; CURRENT APPLICATION DATA:  
; APPLICATION NUMBER: US/08/856,253  
; FILING DATE: Concurrently Herewith  
; CLASSIFICATION: 514  
; PRIOR APPLICATION DATA:  
; APPLICATION NUMBER: US 60/017,678  
; FILING DATE: 16-MAY-1996  
; ATTORNEY/AGENT INFORMATION:  
; NAME: Kitchell, Barbara S.  
; REGISTRATION NUMBER: 33,928  
; REFERENCE/DOCKET NUMBER: TAWK:193  
; TELECOMMUNICATION INFORMATION:  
; TELEPHONE: (512) 418-3000  
; TELEFAX: (512) 474-7577  
; INFORMATION FOR SEQ ID NO: 4:  
; SEQUENCE CHARACTERISTICS:  
; LENGTH: 211 amino acids  
; TYPE: amino acid  
; STRANDEDNESS:  
; TOPOLOGY: linear  
US-08-856-253-4

Query Match 100.0%; Score 1122; DB 3; Length 211;  
Best Local Similarity 100.0%; Pred. No. 4.7e-94;  
Matches 211; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 MRGSHHHHGGDDKVAITISGNKSTNVTHKSEAGTSSVFYKTDGMLPDTTHVRWFL 60  
Db 1 MRGSHHHHGGDDKVAITISGNKSTNVTHKSEAGTSSVFYKTDGMLPDTTHVRWFL 60  
QY 61 NINNEKSYVSKDITIKDQIQGGQQLDLSTLNINVTGTHSNYISGQSAITDFEKAFFGSKI 120  
Db 61 NINNEKSYVSKDITIKDQIQGGQQLDLSTLNINVTGTHSNYISGQSAITDFEKAFFGSKI 120  
QY 121 TVDNTKNTIDVTIPOGYGYSNFSINYKTI NEQOKEFVNNQSAWYQHBGKEEVNGKSF 180  
Db 121 TVDNTKNTIDVTIPOGYGYSNFSINYKTI NEQOKEFVNNQSAWYQHBGKEEVNGKSF 180  
QY 181 NHTVHNINANAGIEGTGKELKVLKODKDTK 211  
Db 181 NHTVHNINANAGIEGTGKELKVLKODKDTK 211

## RESULT 2

US-08-856-253-6  
; Sequence 6, Application US/08856253  
; Patent No. 6288214  
; GENERAL INFORMATION:  
; APPLICANT: Hook, Magnus  
; APPLICANT: Patti, Joseph M.  
; APPLICANT: House-Pompeo, Karen  
; APPLICANT: Symanam, Narayana  
; APPLICANT: Symersky, Jindrich  
; TITLE OF INVENTION: COLLAGEN BINDING PROTEIN COMPOSITIONS  
; AND METHODS OF USE  
; NUMBER OF SEQUENCES: 8  
; CORRESPONDENCE ADDRESS:  
; ADDRESSEE: Arnold, White & Durkee  
; STREET: P.O. Box 4433  
; CITY: Houston  
; STATE: Texas  
; COUNTRY: U.S.  
; ZIP: 77210  
; COMPUTER READABLE FORM:  
; MEDIUM TYPE: Floppy disk  
; COMPUTER: IBM PC compatible  
; OPERATING SYSTEM: PC-DOS/MS-DOS  
; SOFTWARE: Patent In Release #1.0, Version #1.30  
; CURRENT APPLICATION DATA:  
; APPLICATION NUMBER: US/08/856,253  
; FILING DATE: Concurrently Herewith  
; CLASSIFICATION: 514  
; PRIOR APPLICATION DATA:  
; APPLICATION NUMBER: US 60/017,678  
; FILING DATE: 16-MAY-1996  
; ATTORNEY/AGENT INFORMATION:  
; NAME: Kitchell, Barbara S.  
; REGISTRATION NUMBER: 33,928  
; REFERENCE/DOCKET NUMBER: TAMK:193  
; TELECOMMUNICATION INFORMATION:  
; TELEPHONE: (512) 418-3000  
; TELEFAX: (512) 474-7577  
; INFORMATION FOR SEQ ID NO: 6:  
; SEQUENCE CHARACTERISTICS:  
; LENGTH: 512 amino acids  
; TYPE: amino acid  
; STRANDEDNESS:  
; TOPOLOGY: linear  
US-08-856-253-6

Query Match 94.0%; Score 1054.5; DB 3; Length 512;  
Best Local Similarity 64.7%; Pred. No. 2 1e-87;  
Matches 211; Conservative 0; Mismatches 0; Indels 115; Gaps 1;  
QY 1 MRGSHHHHGG-----  
Db 1 MRGSHHHHGGARDISSTNVTDLTVSPSKI EDGKTKTVKMTFDDKNGKI QNGDMIKVAV 60

QY 12 ----- 11  
Db 61 PTSGTVKIEGYKTVPLTVKGEQGVAVITPDGATITFNDKVEKLSVSGFAEPVQGRN 120  
QY 12 -----SDKVAITISGNKSTNVTHKSEAGTSSVFYKTDGMLPDTTHVRWFLINNE 65  
Db 121 LTQNTSDKVAITISGNKSTNVTHKSEAGTSSVFYKTDGMLPDTTHVRWFLINNE 180  
QY 66 KSYVSKDITIKDQIQGGQQLDLSTLNINVTGTHSNYISGQSAITDFEKAFFGSKI TVDNT 125  
Db 181 KSYVSKDITIKDQIQGGQQLDLSTLNINVTGTHSNYISGQSAITDFEKAFFGSKI TVDNT 240  
QY 126 KNTIDVTIPOGYGYSNFSINYKTI NEQOKEFVNNQSAWYQHBGKEEVNGKSFNHTVH 185  
Db 241 KNTIDVTIPOGYGYSNFSINYKTI NEQOKEFVNNQSAWYQHBGKEEVNGKSFNHTVH 300  
QY 186 NINANAGIEGTGKELKVLKODKDTK 211  
Db 301 NINANAGIEGTGKELKVLKODKDTK 326

## RESULT 3

US-08-447-031A-2  
; Sequence 2, Application US/08447031A  
; Patent No. 5851794  
; GENERAL INFORMATION:  
; APPLICANT: GUSS, Bengt  
; APPLICANT: HOOK, Magnus  
; APPLICANT: JONSSON, Hans  
; APPLICANT: LINDBERG, Martin  
; APPLICANT: PATTI, Joseph  
; APPLICANT: SIGNAS, Christer  
; APPLICANT: SWITALSKI, Lech  
; TITLE OF INVENTION: A COLLAGEN BINDING PROTEIN AS WELL AS  
; ITS PREPARATION  
; NUMBER OF SEQUENCES: 8  
; CORRESPONDENCE ADDRESS:  
; ADDRESSEE: Burns, Doane, Swecker & Mathis  
; STREET: P.O. Box 1404  
; CITY: Alexandria  
; STATE: Virginia  
; COUNTRY: United States  
; ZIP: 22313-1404  
; COMPUTER READABLE FORM:  
; MEDIUM TYPE: Floppy disk  
; COMPUTER: IBM PC compatible  
; OPERATING SYSTEM: PC-DOS/MS-DOS  
; SOFTWARE: Patent In Release #1.0, Version #1.30  
; CURRENT APPLICATION DATA:  
; APPLICATION NUMBER: US/08/447,031A  
; FILING DATE: 22-MAY-1995  
; CLASSIFICATION: 435  
; PRIOR APPLICATION DATA:  
; APPLICATION NUMBER: US 07/861,804  
; FILING DATE: 21-AUG-1992  
; PRIOR APPLICATION DATA:  
; APPLICATION NUMBER: WO PCT/SE91/00707  
; FILING DATE: 22-OCT-1991  
; PRIOR APPLICATION DATA:  
; APPLICATION NUMBER: SE 9003374-7  
; FILING DATE: 22-OCT-1990  
; ATTORNEY/AGENT INFORMATION:  
; NAME: McGowan, Malcolm K.  
; REGISTRATION NUMBER: 39,300  
; REFERENCE/DOCKET NUMBER: 012889-006  
; TELECOMMUNICATION INFORMATION:  
; TELEPHONE: (703) 836-6620  
; TELEFAX: (703) 836-2021  
; INFORMATION FOR SEQ ID NO: 2:  
; SEQUENCE CHARACTERISTICS:  
; LENGTH: 1183 amino acids  
; TYPE: amino acid  
; TOPOLOGY: linear

MOLECULE TYPE: protein  
US-08-447-031A-2

Query Match 93.4%; Score 1048; DB 2; Length 1183;  
Best Local Similarity 100.0%; Pred. No. 2.6e-86;  
Matches 200; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 12 SDDKVAITSGNKSNTVTVHKSEAGTSSVFYKGTGMDLPEDTTHVRFNLINNEKSYVSK 71  
DB 144 SDDKVAITSGNKSNTVTVHKSEAGTSSVFYKGTGMDLPEDTTHVRFNLINNEKSYVSK 203

QY 72 DITIKDQIQGGQQLDLSTLNINVTGTHSNYSGOSAITDFEAPGSKITVDNTKTIDV 131  
DB 204 DITIKDQIQGGQQLDLSTLNINVTGTHSNYSGOSAITDFEAPGSKITVDNTKTIDV 263

QY 132 TIPOGYSYNSFSINNYKTKITNEQKEFVNNSQAWYQEHGKBEVNGKSFNHTVHNANANA 191  
DB 264 TIPOGYSYNSFSINNYKTKITNEQKEFVNNSQAWYQEHGKBEVNGKSFNHTVHNANANA 323

QY 192 GIEGTGKELKVLKQDKDK 211  
DB 324 GIEGTGKELKVLKQDKDK 343

RESULT 4  
US-08-856-253-2  
Sequence 2, Application US/08856253  
Patent No. 6288214  
GENERAL INFORMATION:  
APPLICANT: Hook, Magnus  
APPLICANT: Patti, Joseph M.  
APPLICANT: House-Pompeo, Karen  
APPLICANT: Sthanam, Narayana  
APPLICANT: Symersky, Jindrich  
TITLE OF INVENTION: COLLAGEN BINDING PROTEIN COMPOSITIONS  
TITLE OF INVENTION: AND METHODS OF USE  
NUMBER OF SEQUENCES: 8  
CORRESPONDENCE ADDRESS:  
ADDRESSEE: Arnold, White & Durkee  
STREET: P.O. Box 4433  
CITY: Houston  
STATE: Texas  
COUNTRY: U.S.  
ZIP: 77210  
COMPUTER READABLE FORM:  
MEDIUM TYPE: Floppy disk  
COMPUTER: IBM PC compatible  
OPERATING SYSTEM: PC-DOS/MS-DOS  
SOFTWARE: Patentin Release #1.0, Version #1.30  
CURRENT APPLICATION DATA:  
APPLICATION NUMBER: US/08/856,253  
FILING DATE: Concurrently Herewith  
CLASSIFICATION: 514  
PRIOR APPLICATION DATA:  
APPLICATION NUMBER: US 60/017,678  
FILING DATE: 16-MAY-1996  
ATTORNEY/AGENT INFORMATION:  
NAME: Kitchell, Barbara S.  
REGISTRATION NUMBER: 33,928  
REFERENCE/DOCKET NUMBER: TAMK:193  
TELECOMMUNICATION INFORMATION:  
TELEPHONE: (512) 418-3000  
TELEFAX: (512) 474-7577  
INFORMATION FOR SEQ ID NO: 2:  
SEQUENCE CHARACTERISTICS:  
LENGTH: 159 amino acids  
TYPE: amino acid  
STRANDEDNESS:  
TOPOLOGY: linear  
US-08-856-253-2

Query Match 74.0%; Score 830; DB 3; Length 159;  
Best Local Similarity 96.4%; Pred. No. 8.9e-68;

Matches 159; Conservative 0; Mismatches 0; Indels 6; Gaps 1;

QY 1 MRGSHHHHGGSDDKVATITSGNKSNTVTVHKSEAGTSSVFYKGTGMDLPEDTTHVRFNL 60  
DB 1 MRGSHHHHGGSDDKVATITSGNKSNTVTVHKSEAGTSSVFYKGTGMDLPEDTTHVRFNL 54

QY 61 NINNEKSYVSKDITIKDQIQGGQQLDLSTLNINVTGTHSNYSGOSAITDFEAPGSKI 120  
DB 55 NINNEKSYVSKDITIKDQIQGGQQLDLSTLNINVTGTHSNYSGOSAITDFEAPGSKI 114

QY 121 TVDNTKNTIDVTIPGYSYNSFSINNYKTKITNEQKEFVNNSQA 165  
DB 115 TVDNTKNTIDVTIPGYSYNSFSINNYKTKITNEQKEFVNNSQA 159

RESULT 5  
US-09-107-532A-6999  
Sequence 6999, Application US/09107532A  
Patent No. 6583275  
GENERAL INFORMATION:  
APPLICANT: Lynn A Doucette-Stamm and David Bush  
TITLE OF INVENTION: NUCLEIC ACID AND AMINO ACID SEQUENCES RELATING TO ENTEROCOCCUS FAECIUM FOR DIAGNOSTICS AND THERAPEUTICS  
NUMBER OF SEQUENCES: 7310  
CORRESPONDENCE ADDRESS:  
ADDRESSEE: GENOME THERAPEUTICS CORPORATION  
STREET: 100 Beaver Street  
CITY: Waltham  
STATE: Massachusetts  
COUNTRY: USA  
ZIP: 02354  
COMPUTER READABLE FORM:  
MEDIUM TYPE: CD-ROM ISO9660  
COMPUTER: PC  
OPERATING SYSTEM: <Unknown>  
SOFTWARE: ASCII  
CURRENT APPLICATION DATA:  
APPLICATION NUMBER: US/09/107,532A  
FILING DATE: 30-Jun-1998  
PRIOR APPLICATION DATA:  
APPLICATION NUMBER: 60/085,598  
FILING DATE: 14 May 1998  
APPLICATION NUMBER: 60/051571  
FILING DATE: July 2, 1997  
ATTORNEY/AGENT INFORMATION:  
NAME: Ariniello, Pamela Deneka  
REGISTRATION NUMBER: 40,489  
REFERENCE/DOCKET NUMBER: GTC-012  
TELECOMMUNICATION INFORMATION:  
TELEPHONE: (781)893-5007  
TELEFAX: (781)893-8277  
INFORMATION FOR SEQ ID NO: 6999:  
SEQUENCE CHARACTERISTICS:  
LENGTH: 731 amino acids  
TYPE: amino acid  
TOPOLOGY: linear  
MOLECULE TYPE: protein  
HYPOTHETICAL: YES  
ORIGINAL SOURCE:  
ORGANISM: Enterococcus faecium  
FEATURE:  
NAME/KEY: misc feature  
LOCATION: (B) LOCATION 1...731  
SEQUENCE DESCRIPTION: SEQ ID NO: 6999:  
US-09-107-532A-6999

Query Match 43.8%; Score 491; DB 4; Length 731;  
Best Local Similarity 48.5%; Pred. No. 3.8e-36;  
Matches 95; Conservative 35; Mismatches 64; Indels 2; Gaps 2;

QY 18 TITSGNKSNTVTVHKSEAGTSSVFYKGTGMDLPEDTTHVRFNLINNEKSYVSKDITIK 76  
DB 162 TVTSGDKATVNTVTPASGSSSSVFYKGTGMDLPEDTTHVRFNLINNNNGTYVEQPVKIS 221



; FILING DATE:
; CLASSIFICATION:
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER:
; FILING DATE:
; ATTORNEY/AGENT INFORMATION:
; NAME: A. Anders Brookes
; REGISTRATION NUMBER: 36,373
; REFERENCE/DOCKET NUMBER: PB369P2
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (301) 309-8504
; TELEFAX: (301) 309-8512
; INFORMATION FOR SEQ ID NO: 464:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 688 amino acids
; TYPE: amino acid
; STRANDEDNESS: single
; TOPOLOGY: linear
; MOLECULE TYPE: protein
; US-09-071-035-464

Query Match 10.2%; Score 114.5; DB 4; Length 688;
Best Local Similarity 25.5%; Pred. No. 0.045;
Matches 54; Conservative 36; Mismatches 93; Indels 29; Gaps 9;

QY 12 SDDKVAITSGNKSTNVTVHKS-EAGTSSVFYKTDGMLPEDTTHVRWFLNINNEKSYVS 70
Db 172 SYDNTASTYTNQSSRDVTGKVSIOHGGSV--KKGGEYHKDDPDHVVHVMINGAQS-VL 228

QY 71 KDITIKDQIOGGQQLDLSTLNINVTGTHSNYSGQSAITDFEKAFFGSKITVDNTKN-TI 129
Db 229 DDVITDTPSPNQVLDPSLVI--YGTN-----VTEGTTITPKSVILEEGKDYTL 277

QY 130 DVTIPIQGVGSYN-----SFSINYTKITNEQ--QKEFVNNSQAWYQEHGKEEVNG 177
Db 278 EVTTDNETGQKIVVVKMAHIEAPYMEYRSLVTSAGSTDTVSN-QVSIITGNGSEVVG 336

QY 178 KSFNHTVHNINANAGIEGTGKELKVLKQDKD 209
Db 337 DDNGDVVVDIDHSGGHATGKGKIQLKKTAMD 368

RESULT 9
US-09-071-035-458
; Sequence 458, Application US/09071035
; Patent No. 6448043
; GENERAL INFORMATION:
; APPLICANT: Gil H. Choi
; TITLE OF INVENTION: Enterococcus faecalis Polynucleotides and Polypeptides
; NUMBER OF SEQUENCES: 496
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Human Genome Sciences, Inc.
; STREET: 9410 Key West Avenue
; CITY: Rockville
; STATE: Maryland
; COUNTRY: USA
; ZIP: 20850
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Diskette, 3.50 inch, 1.4Mb storage
; COMPUTER: HP Vectra 486/33
; OPERATING SYSTEM: MSDOS version 6.2
; SOFTWARE: ASCII Text
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/09/071,035
; FILING DATE:
; CLASSIFICATION:
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER:
; FILING DATE:
; ATTORNEY/AGENT INFORMATION:
; NAME: A. Anders Brookes
; REGISTRATION NUMBER: 36,373
; REFERENCE/DOCKET NUMBER: PB369P2
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (301) 309-8504
; TELEFAX: (301) 309-8512
; INFORMATION FOR SEQ ID NO: 462:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 2032 amino acids
; TYPE: amino acid
; STRANDEDNESS: single
; TOPOLOGY: linear

; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (301) 309-8504
; TELEFAX: (301) 309-8512
; INFORMATION FOR SEQ ID NO: 458:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 2032 amino acids
; TYPE: amino acid
; STRANDEDNESS: single
; TOPOLOGY: linear
; MOLECULE TYPE: protein
; US-09-071-035-458

Query Match 10.2%; Score 114.5; DB 4; Length 2032;
Best Local Similarity 25.5%; Pred. No. 0.2;
Matches 54; Conservative 36; Mismatches 93; Indels 29; Gaps 9;

QY 12 SDDKVAITSGNKSTNVTVHKS-EAGTSSVFYKTDGMLPEDTTHVRWFLNINNEKSYVS 70
Db 908 SYDNTASTYTNQSSRDVTGKVSIOHGGSV--KKGGEYHKDDPDHVVHVMINGAQS-VL 964

QY 71 KDITIKDQIOGGQQLDLSTLNINVTGTHSNYSGQSAITDFEKAFFGSKITVDNTKN-TI 129
Db 965 DDVITDTPSPNQVLDPSLVI--YGTN-----VTEGTTITPKSVILEEGKDYTL 1013

QY 130 DVTIPIQGVGSYN-----SFSINYTKITNEQ--QKEFVNNSQAWYQEHGKEEVNG 177
Db 1014 EVTTDNETGQKIVVVKMAHIEAPYMEYRSLVTSAGSTDTVSN-QVSIITGNGSEVVG 1072

QY 178 KSFNHTVHNINANAGIEGTGKELKVLKQDKD 209
Db 1073 DDNGDVVVDIDHSGGHATGKGKIQLKKTAMD 1104

RESULT 10
US-09-071-035-462
; Sequence 462, Application US/09071035
; Patent No. 6448043
; GENERAL INFORMATION:
; APPLICANT: Gil H. Choi
; TITLE OF INVENTION: Enterococcus faecalis Polynucleotides and Polypeptides
; NUMBER OF SEQUENCES: 496
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Human Genome Sciences, Inc.
; STREET: 9410 Key West Avenue
; CITY: Rockville
; STATE: Maryland
; COUNTRY: USA
; ZIP: 20850
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Diskette, 3.50 inch, 1.4Mb storage
; COMPUTER: HP Vectra 486/33
; OPERATING SYSTEM: MSDOS version 6.2
; SOFTWARE: ASCII Text
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/09/071,035
; FILING DATE:
; CLASSIFICATION:
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER:
; FILING DATE:
; ATTORNEY/AGENT INFORMATION:
; NAME: A. Anders Brookes
; REGISTRATION NUMBER: 36,373
; REFERENCE/DOCKET NUMBER: PB369P2
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (301) 309-8504
; TELEFAX: (301) 309-8512
; INFORMATION FOR SEQ ID NO: 462:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 2032 amino acids
; TYPE: amino acid
; STRANDEDNESS: single
; TOPOLOGY: linear

```
; MOLECULE TYPE: protein
US-09-071-035-462

Query Match      10.2%; Score 114.5; DB 4; Length 2032;
Best Local Similarity 25.5%; Pred. No. 0.2;
Matches 54; Conservative 36; Mismatches 93; Indels 29; Gaps 9;

QY 12 SDDKVATITSGNKSTNVTVHKS--EAGTSSVFYKGTGDMLPEDTTHVRWFLNINNEKSYVS 70
DB 908 SYDNTASYTNOGSSRDVTGKVSIOHGGSV--KKGGEYHKDDPDHVVYHVMINGAQS-VL 964
QY 71 KDIITKDOIQQGQQLDLSTLNINVTGTHSNYYSQSAITDFEKAFFPGSKITVDNTKN-TI 129
DB 965 DDVWITDTPSPNQVLDPESLVI--YGTN-----VTEGTTTPDKSVILEEGKDYTL 1013
QY 130 DVTIPQGYGSYN-----SFSINYKTKITNEQ--QKEFVNNSQAWYQEHGKEEVNG 177
DB 1014 EVTTDNETGQOKIVVVKMAHIEAPYMEYRSLVTSAAAGSTDVSN-QVSIITGNGSEVVHG 1072
QY 178 KSFNHTVHNINANAGIEGTGKELVKLQDKD 209
DB 1073 DDNGDVVDIDHSGGHATGKGIQLKKTAMD 1104

RESULT 12
US-09-134-000C-6612
; Sequence 6612, Application US/09134000C
; Patent No. 6617156
; GENERAL INFORMATION:
; APPLICANT: Lynn Doucette-Stamm et al
; TITLE OF INVENTION: NUCLEIC ACID AND AMINO ACID SEQUENCES RELATING TO
; FILE REFERENCE: ENTEROCOCCUS FAECALIS FOR DIAGNOSTICS AND THERAPEUTICS
; CURRENT APPLICATION NUMBER: US/09/134.000C
; CURRENT FILING DATE: 1998-08-13
; PRIOR APPLICATION NUMBER: US 60/055,778
; PRIOR FILING DATE: 1997-08-15
; NUMBER OF SEQ ID NOS: 6812
; SOFTWARE: Patent in version 3.1
; SEQ ID NO 6612
; LENGTH: 2054
; TYPE: PRT
; ORGANISM: Enterococcus faecalis
US-09-134-000C-6612

Query Match      10.2%; Score 114.5; DB 4; Length 2054;
Best Local Similarity 25.5%; Pred. No. 0.21;
Matches 54; Conservative 36; Mismatches 93; Indels 29; Gaps 9;

QY 12 SDDKVATITSGNKSTNVTVHKS--EAGTSSVFYKGTGDMLPEDTTHVRWFLNINNEKSYVS 70
DB 930 SYDNTASYTNOGSSRDVTGKVSIOHGGSV--KKGGEYHKDDPDHVVYHVMINGAQS-VL 986
QY 71 KDIITKDOIQQGQQLDLSTLNINVTGTHSNYYSQSAITDFEKAFFPGSKITVDNTKN-TI 129
DB 987 DDVWITDTPSPNQVLDPESLVI--YGTN-----VTEGTTTPDKSVILEEGKDYTL 1035
QY 130 DVTIPQGYGSYN-----SFSINYKTKITNEQ--QKEFVNNSQAWYQEHGKEEVNG 177
DB 1036 EVTTDNETGQOKIVVVKMAHIEAPYMEYRSLVTSAAAGSTDVSN-QVSIITGNGSEVVHG 1094
QY 178 KSFNHTVHNINANAGIEGTGKELVKLQDKD 209
DB 1095 DDNGDVVDIDHSGGHATGKGIQLKKTAMD 1126

RESULT 13
US-09-023-082A-30
; Sequence 30, Application US/09023082A
; Patent No. 6077692
; GENERAL INFORMATION:
; APPLICANT: RUBEN, STEVEN M.
; APPLICANT: JIMENEZ, PABLO
; APPLICANT: DUAN, D. ROXANNE
; APPLICANT: RAMPY, MARK A.
; APPLICANT: MENDRICK, DONNA
; APPLICANT: ZHANG, JUN
; APPLICANT: NI, JIAN
; APPLICANT: MOORE, PAUL A.
; APPLICANT: COLEMAN, TIMOTHY A.
; APPLICANT: GRUBER, JOACHIM R.
; APPLICANT: DILLON, PATRICK J.

; MOLECULE TYPE: protein
US-09-071-035-466

Query Match      10.2%; Score 114.5; DB 4; Length 2032;
Best Local Similarity 25.5%; Pred. No. 0.2;
Matches 54; Conservative 36; Mismatches 93; Indels 29; Gaps 9;

QY 12 SDDKVATITSGNKSTNVTVHKS--EAGTSSVFYKGTGDMLPEDTTHVRWFLNINNEKSYVS 70
DB 908 SYDNTASYTNOGSSRDVTGKVSIOHGGSV--KKGGEYHKDDPDHVVYHVMINGAQS-VL 964
QY 71 KDIITKDOIQQGQQLDLSTLNINVTGTHSNYYSQSAITDFEKAFFPGSKITVDNTKN-TI 129
DB 965 DDVWITDTPSPNQVLDPESLVI--YGTN-----VTEGTTTPDKSVILEEGKDYTL 1013
QY 130 DVTIPQGYGSYN-----SFSINYKTKITNEQ--QKEFVNNSQAWYQEHGKEEVNG 177
DB 1014 EVTTDNETGQOKIVVVKMAHIEAPYMEYRSLVTSAAAGSTDVSN-QVSIITGNGSEVVHG 1072
QY 178 KSFNHTVHNINANAGIEGTGKELVKLQDKD 209
DB 1073 DDNGDVVDIDHSGGHATGKGIQLKKTAMD 1104

RESULT 11
US-09-071-035-466
; Sequence 466, Application US/09071035
; Patent No. 6448043
; GENERAL INFORMATION:
; APPLICANT: Gil H. Choi
; TITLE OF INVENTION: Enterococcus faecalis Polynucleotides and Polypeptides
; NUMBER OF SEQUENCES: 496
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Human Genome Sciences, Inc.
; STREET: 9410 Key West Avenue
; CITY: Rockville
; STATE: Maryland
; COUNTRY: USA
; ZIP: 20850
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Diskette, 3.50 inch, 1.4Mb storage
; COMPUTER: HP Vectra 486/33
; OPERATING SYSTEM: MSDOS version 6.2
; SOFTWARE: ASCII Text
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/09/071,035
; FILING DATE:
; CLASSIFICATION:
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER:
; FILING DATE:
; ATTORNEY/AGENT INFORMATION:
; NAME: A. Anders Brookes
; REGISTRATION NUMBER: 36,373
; REFERENCE/DOCKET NUMBER: PB369P2
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (301) 309-8504
; TELEFAX: (301) 309-8512
; INFORMATION FOR SEQ ID NO: 466:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 2032 amino acids
; TYPE: amino acid
; STRANDEDNESS: single
; TOPOLOGY: linear
; MOLECULE TYPE: protein
US-09-071-035-466

Query Match      10.2%; Score 114.5; DB 4; Length 2032;
Best Local Similarity 25.5%; Pred. No. 0.2;
Matches 54; Conservative 36; Mismatches 93; Indels 29; Gaps 9;

QY 12 SDDKVATITSGNKSTNVTVHKS--EAGTSSVFYKGTGDMLPEDTTHVRWFLNINNEKSYVS 70
```



```

Qy      1 MRGSHHHHGGDDKVIATITSGNKSTNTVHKSEAGTSSVFYKYTKDMLPEDTTTHRWFL 60
         |||||:|||||:|||||:|||||:|||||:|||||:|||||:|||||:|||||:
Db      1 MRGSHHHHGGSCALGDQMVSPEATNSSSSSFSPSAGRHVRSVNHLCQD---VRWRK 57
         :|||||:|||||:|||||:|||||:|||||:|||||:|||||:|||||:
Qy     61 NINNEKSVSKDIITIKOIQGGQQ-----LDLSTLNINVGT---HSNYVGOSAITD 110
         :|||:|||||:|||||:|||||:|||||:|||||:|||||:|||||:
Db     58 LPSFTKYPLK--IEKNGVSGTKKENCPCYSILEITSVEIGVVAVKAINSYY---LAMNK 112
         :|||:|||||:|||||:|||||:|||||:|||||:|||||:|||||:
Qy    111 FEKAFPGSKITVD-NTRKNTIDVTIPOGYSYNFSFNINYKTITNBQOKKEFVNNSOAQYOE 169
         :|||:|||||:|||||:|||||:|||||:|||||:|||||:|||||:
Db    113 KGKLVGSKFEFNNDCKLERIE--ENGINTYASFN-----NQOH 147
         :|||:|||||:|||||:|||||:|||||:|||||:|||||:|||||:
Qy    170 HGKE----EVNGK 178
         :|||:|||||:|||||:|||||:|||||:|||||:|||||:|||||:

```

APPLICATION NUMBER: US/08/728,470  
FILING DATE:  
CLASSIFICATION: 424  
PRIOR APPLICATION NUMBER: US 08/302,832  
FILING DATE: 16-MAR-1993  
PRIOR APPLICATION DATA:  
APPLICATION NUMBER: US PCT/US93/02166  
FILING DATE: 16-MAR-1993  
PRIOR APPLICATION DATA:  
APPLICATION NUMBER: GB 9205704.1  
FILING DATE: 16-MAR-1992  
ATTORNEY/AGENT INFORMATION:  
NAME: Berkatresser, Jerry W  
REGISTRATION NUMBER: 22,651  
REFERENCE/DOCKET NUMBER: 1038-633  
TELECOMMUNICATION INFORMATION:  
TELEPHONE: (703) 415-0810  
TELEFAX: (703) 415-0813  
INFORMATION FOR SEQ ID NO: 9:  
SEQUENCE CHARACTERISTICS:  
LENGTH: 1338 amino acids  
TYPE: amino acid  
STRANDEDNESS: single  
TOPOLOGY: linear  
US-08-728-470-9

Query Match 9.8%; Score 110.5; DB 2; Length 1338;  
Best Local Similarity 25.5%; Pred. No. 0.26;  
Matches 61; Conservative 28; Mismatches 89; Indels 61; Gaps 10;

QY 12 SDKVATITSGNKST-----NVTVKSEA-----GTSSVFFYKTDMLPETHVRFILN 61  
DB 694 SDDTYRTIKGNISKSGDLNIDKKSDAEIQIGNIS---QKEGN-----LT 738

QY 62 INNEKSYVSKDITIKDQIQGGQQLDLSTLINVNT-GTHSNYYSGQSAITDFEKAFFGSKI 120  
DB 739 ISSDKVNITNQITIKAGVEGGRSDSEAEANLTIQTKELKLAGDLNISGFNKA----EI 794

QY 121 TVDNTYNTIDVTIPOGYGSY-----NSFSINYTKITNEQQKEFV 160  
DB 795 T---AKNGSDLTIGNASGGNADAKKVTFDKVKDSKISTDGHNVTLNSEVKTSGSSNAGN 851

QY 161 NNSQAWYQEHGKEEVNGKSFNHTVHNINANAG----IEGT-----VKGELKVLKQDKDK 211  
DB 852 DNSTGLTISAKDVTNNVNTSHKTNISAAAGNVTTKETGINATGTSVEVTAQNGTIK 910

Search completed: July 27, 2004, 10:10:29  
Job time : 20 secs

GenCore version 5.1.6  
Copyright (c) 1993 - 2004 CompuGen Ltd.

OM protein - protein search, using sw model  
Run on: July 27, 2004, 10:06:36 ; Search time 16 Seconds  
(without alignments)  
1268.526 Million cell updates/sec

Title: US-09-813-820-4  
Perfect score: 1122  
Sequence: 1 MRGSHHHHHGSDDKVATIT.....GIEGTVKGELKVLKQDKDTK 211

Scoring table: BLOSUM62  
Gapop 10.0 , Gapext 0.5

Searched: 283366 seqs, 96191526 residues  
Total number of hits satisfying chosen parameters: 283366

Minimum DB seq length: 0  
Maximum DB seq length: 2000000000  
Post-processing: Minimum Match 0%  
Maximum Match 100%  
Listing first 45 summaries

Database : PIR 78:\*  
1: pir1:\*  
2: pir2:\*  
3: pir3:\*  
4: pir4:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	ID	Description
1	911	81.2	1185	2 A42404	collagen adhesin -
2	194	17.3	37	2 A48620	adhesin - Staphylo
3	131	11.7	1092	2 S42798	fibronectin-bindin
4	119	10.6	336	2 D86710	collagen adhesin l
5	101.5	9.0	542	2 A98021	hypothetical prote
6	100	8.9	1530	2 A41396	peptidoglycan anch
7	100	8.9	1536	2 A43855	high-molecular-wei
8	99.5	8.9	266	2 D97203	hypothetical prote
9	99.5	8.9	463	2 AG1542	wall associated pr
10	99.5	8.9	592	2 A81888	probable surface f
11	98.5	8.8	4919	2 T31105	hypothetical prote
12	98	8.7	1117	2 S33851	fibronectin-bindin
13	98	8.7	1248	2 C89874	autolysin (importe
14	96.5	8.6	848	2 C70203	DNA topoisomerase
15	96	8.6	392	2 A53580	neurexin III beta
16	96	8.6	704	1 S39004	finger protein MSN
17	95	8.5	359	2 E86706	hypothetical prote
18	95	8.5	666	2 A42296	lysosome 2 (EC 3.2
19	95	8.5	2020	2 C48399	ABC-type transport
20	94.5	8.4	188	2 E90065	hypothetical prote
21	94.5	8.4	2902	2 C71953	toxin-like outer m
22	94.5	8.4	3890	2 C89921	hypothetical prote
23	94	8.4	2380	2 E71604	hypothetical prote
24	93.5	8.3	532	2 T06029	hypothetical prote
25	93.5	8.3	1237	2 S39791	neurotoxin - Clost
26	93.5	8.3	2340	2 B71704	cell surface antig
27	92.5	8.2	681	2 AC1458	surface anchored p
28	92.5	8.2	711	2 S73898	DNA topoisomerase
29	92.5	8.2	1943	2 B64596	toxin-like outer m

30 92.5 8.2 2910 2 T28156 DNA-directed RNA p  
31 92 8.2 1037 2 T13350 transcription fact  
32 92 8.2 1983 2 G86643 hypothetical prote  
33 92 8.2 2500 2 G71609 hypothetical prote  
34 91.5 8.2 261 2 S16619 opacity protein op  
35 91.5 8.2 622 2 AF0169 probable exported  
36 91.5 8.2 2150 2 S71629 sensory transducti  
37 91.5 8.2 2271 2 F90073 hypothetical prote  
38 91 8.1 452 2 AE1710 hypothetical prote  
39 91 8.1 463 2 A44808 cellulase (EC 3.2.  
40 91 8.1 471 2 H97200 ferrichrome-bindin  
41 91 8.1 578 2 B90578 lipoprotein a (limp  
42 91 8.1 953 2 C89824 hypothetical prote  
43 91 8.1 2269 2 T28677 lipoprotein B-1  
44 90.5 8.1 784 2 JH0101 apolipoprotein B-1  
45 90 8.0 669 2 S14535 asparagine-rich pr

ALIGNMENTS

RESULT 1  
A42404  
collagen adhesin - Staphylococcus aureus  
C:Species: Staphylococcus aureus  
C:Date: 04-Mar-1993 #sequence\_revision 18-Nov-1994 #text\_change 10-Nov-1995  
C:Accession: A42404; S27665  
R:Patti, J.M.; Jonsson, H.; Guss, B.; Switalski, L.M.; Wiberg, K.; Lindberg, M.; Hook, J.  
J. Biol. Chem. 267, 4766-4772, 1992  
A:Title: Molecular characterization and expression of a gene encoding a Staphylococcus  
A:Reference number: A42404; MUID:92165839; PMID:1311320  
A:Contents: FDA 574  
A:Accession: A42404  
A>Status: preliminary  
A:Molecule type: DNA  
A:Residues: 1-1185 <PAT>  
A:Cross-references: EMBL:N81736  
A>Note: sequence extracted from NCBI backbone (NCBIP:83982)

Query Match 81.2%; Score 911; DB 2; Length 1185;  
Best Local Similarity 90.1%; Pred. No. 76-58;  
Matches 182; Conservative 2; Mismatches 16; Indels 2; Gaps 2;  
QY 12 SDDKVATITSGNKSTNVTVHKSEAGTSSVFYY-KTGDMLPBDTTHVRFLNNKESYV 69  
DB 144 SDDKVATITSGNKSTNVTGKREPVVFLINKSGKICYQEDTTHVRFLNNKESYV 203  
QY 70 SKDITIKDQIQGGQQLDLSTLNINVTGTHSNYYSGSAITDFEKAPPGSKIIVDNTKNTI 129  
DB 204 SKDITIKDQIQGGQQLDLSTLNINVTGTHSNYYSGSAITDFEKAPPGSKIIVDNTKNTI 263  
QY 130 DVTIPQGVSYNSPFSINVKYKITNEQKEFVNNSQAWYQEHGKEEVNGKSFNHTVHNINA 189  
DB 264 DVTIPQGVSYNSPFSINVKYKITNEQKEFVNNSQAWYQEHGKEEVNGKSFNHTVHNINA 323  
QY 190 NAGIEGTVKGELKVLKQDKDTK 211  
DB 324 NAGIEGTVKGELKVLKQDKDTK 345

RESULT 2  
A48620  
adhesin - Staphylococcus aureus (fragment)  
C:Species: Staphylococcus aureus  
C:Date: 07-Apr-1994 #sequence\_revision 18-Nov-1994 #text\_change 03-Mar-1995  
C:Accession: A48620  
R:Patti, J.M.; Boles, J.O.; Hook, M.  
Biochemistry 32, 11428-11435, 1993  
A:Title: Identification and biochemical characterization of the ligand binding domain of  
A:Reference number: A48620; MUID:94032261; PMID:8218209  
A:Contents: FDA 574  
A:Accession: A48620  
A>Status: preliminary; not compared with conceptual translation

A:Molecule type: nucleic acid; protein  
A:Residues: 1-37 <PAT>  
A>Note: sequence extracted from NCBI backbone (NCBIP:138726)

Query Match 17.3%; Score 194; DB 2; Length 37;  
Best Local Similarity 100.0%; Pred. No. 5.2e-08;  
Matches 37; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 19 ITSNGKSTNVTVHKSEAGTSSVFYKGTGMDLPEDTTH 55  
DB 1 ITSNGKSTNVTVHKSEAGTSSVFYKGTGMDLPEDTTH 37

RESULT 3  
S42798  
fibronectin-binding protein - Streptococcus "equisimilis"  
C:Species: Streptococcus "equisimilis"  
C:Date: 06-Jan-1995 #sequence\_revision 06-Jan-1995 #text\_change 15-Oct-1999  
C:Accession: S42798  
R:Signaes, C.  
submitted to the EMBL Data Library, December 1993  
A:Reference number: S42798  
A:Accession: S42798  
A>Status: preliminary  
A:Molecule type: DNA  
A:Residues: 1-1092 <SIG>  
A:Cross-references: EMBL:229088; NID:g436135; PIDN:CAA82330.1; PID:g436136  
C:Keywords: fibronectin binding

Query Match 11.7%; Score 131; DB 2; Length 1092;  
Best Local Similarity 26.5%; Pred. No. 0.13;  
Matches 58; Conservative 31; Mismatches 88; Indels 42; Gaps 13;

QY 19 ITSNGKSTNVTVHKSEAGTSS-VFYKGTGMDLPEDTTH---VRWFLNIN-NEKSVVSXDI 73  
DB 152 VISGTHAEVWQKRGEGGTATDNIIYKSGDI--PDTQEDSVRWSFTFNAARKSTNGVGF 209

QY 74 TTIKDOIQQGQQLD-----STLNINVTGTHSNVYSGQSAITDPE-----KAPPGSKITVD 123  
DB 210 LVTDILDSTNWDVKNLREKAIINFTGG-----WIGDELLKSKNWLKSEAKENYGIKVE 265

QY 124 NTKNTIDVTIPO-----GYGSY-----NSFSINYKTKITNEQOKEFVNNSQAWY-QEHGKE 173  
DB 266 FSGQIVSINPEKIQVDNGSYQELNESEMQHLYAKI-----KKEVLNSSIEYVRESKV 321

QY 174 EVNGKSF-----NHT-----VHNINANAGIBOTVKGELKVLK 205  
DB 322 EVRGENWIPDPNSTSAYVQILRQGGWAKGTVRGEVRLK 360

RESULT 4  
D86710  
collagen adhesin [imported] - Lactococcus lactis subsp. lactis (strain IL1403)  
C:Species: Lactococcus lactis subsp. lactis  
C:Date: 23-Mar-2001 #sequence\_revision 23-Mar-2001 #text\_change 03-Aug-2001  
C:Accession: D86710  
R:Boletín, A.; Wincker, P.; Mauger, S.; Jaillon, O.; Malarne, K.; Weissenbach, J.; Ehrlich  
Genome Res. 11, 731-753, 2001  
A:Title: The complete genome sequence of the lactic acid bacterium Lactococcus lactis se  
A:Reference number: A86625; MUID:21235186; PMID:11337471  
A:Accession: D86710  
A:Molecule type: DNA  
A>Status: preliminary  
A:Residues: 1-336 <STO>  
A:Cross-references: GB:AE005176; PID:gn12723592; PIDN:AAK04782.1; GSPDB:GN00146  
A:Experimental source: strain IL1403  
C:Genetics:  
A:Gene: icaC

Query Match 10.6%; Score 119; DB 2; Length 336;  
Best Local Similarity 25.2%; Pred. No. 0.22;  
Matches 52; Conservative 28; Mismatches 68; Indels 50; Gaps 11;

QY 24 KSTNVTVHKSEAGTSSV-----FYKGTGMDLPEDTTHVRWFLNINNEKSYVSK 71  
DB 71 EDTRVVDWKDGGTTEVNI GPATGDKDEVLYKKGWDENDSTLIHQVRINYAKENTQK 130

QY 72 DITIKDOIQQGQQLD---DLSTLNINVTGTHSNY-----YSGQSAITDPEKAPPGSKITVDN 124  
DB 131 AI-VTDIIGGNQNLVSGSISVANVTYSSDGENVVDVSYYPQASI--LENGVNGFTVNLGD 187

QY 125 TKNITDVTIPQGYGSYNSFSINYKTKITN---EQQKEFVNNSQAWYQEHGKEEVNGKSPN 181  
DB 188 ISNTI-----IIDYSTRATDGLSQO-----YENRG--ELTGENIE 221

QY 182 -----HTVHN-INANAGIEGTGKE 200  
DB 222 KQVVEVHTPNNGNGNASKMLISGE 247

RESULT 5  
A98021  
hypothetical protein appA [imported] - Streptococcus pneumoniae (strain R6)  
C:Species: Streptococcus pneumoniae  
C:Date: 22-Oct-2001 #sequence\_revision 22-Oct-2001 #text\_change 22-Oct-2001  
C:Accession: A98021  
R:Hoskins, J.A.; Alborn Jr., W.; Arnold, J.; Blaszcak, L.; Burgett, S.; DeHoff, B.S.; E  
e, R.; LeBlanc, D.J.; Lee, L.N.; Lefkowitz, E.J.; Lu, J.; Mataushima, P.; McAhren, S.; M  
y, P.; Sun, P.M.; Winkler, M.E.  
J. Bacteriol. 183, 5709-5717, 2001  
A:Authors: Yang, Y.; Young-Bellido, M.; Zhao, G.; Zook, C.; Baltz, R.H.; Jaskunas, S.R.;  
A:Title: Genome of the Bacterium Streptococcus pneumoniae Strain R6.  
A:Reference number: A97872; MUID:21429245; PMID:11544234  
A:Accession: A98021  
A>Status: preliminary  
A:Molecule type: DNA  
A:Residues: 1-542 <KUR>  
A:Cross-references: GB:AE007317; PIDN:AAK99997.1; PID:g15458827; GSPDB:GN00174  
C:Genetics:  
A:Gene: appA

Query Match 9.0%; Score 101.5; DB 2; Length 542;  
Best Local Similarity 25.3%; Pred. No. 7.2;  
Matches 49; Conservative 28; Mismatches 62; Indels 55; Gaps 10;

QY 20 TSGNKSTNVTVHKSEAGTSSVFYK-----TGDMLP----- 50  
DB 25 SSGSKKSDP-----SESGKEKVFYLSGFLQGTGDRIPYQWNGDGLVPVLLYRTLLMADSK 80

QY 51 -EDTTH--VRWFLNINNEKSYVSKDITIKDOIQ--GGQQLDLSTLNINV-----TG 96  
DB 81 YEKSTPDLAKDWKKSDDEKTY---EFTLKGLKWSGDGEALTASDVEFSVKLAKASVING 137

QY 97 THSNYYSQSAITDPE--EKAPPGSKITVDNTKNTIDVTIPOG--YGSVNSFSINYKTKIT 152  
DB 138 IFPSTFTKIEGAKFDKDDKADKISGLTVDGNKVTFKLTAPVGPDFEQMMTQFFILPKHKL 197

QY 153 NEQOKEFVNNSQAW 166  
DB 198 SENPLE-LNNSFW 210

RESULT 6  
AH1396  
peptidoglycan anchored protein (LPXTG motif) [imported] - Listeria monocytogenes (strain  
C:Species: Listeria monocytogenes  
C:Date: 27-Nov-2001 #sequence\_revision 27-Nov-2001 #text\_change 27-Nov-2001  
C:Accession: AH1396  
R:Glaser, P.; Frangoul, L.; Buchrieser, C.; Amend, A.; Baquero, F.; Berche, P.; Bloecker  
.; Dominguez-Bernal, G.; Duchaud, E.; Durand, L.; Dussurget, O.; Entian, K.D.; Fsihi, H.  
D.; Jones, L.M.; Karst, U.  
Science 294, 849-852, 2001  
A:Authors: Kretz, J.; Kuhn, M.; Kunst, F.; Kurapkat, G.; Madueno, E.; Maitournam, A.; Ma  
ok, C.; Schlueter, T.; Simoes, N.; Tierrez, A.; Vazquez-Boland, J.A.; Voss, H.; Wehland,  
A:Title: Comparative genomics of Listeria species.  
A:Reference number: AB1077; MUID:21537279; PMID:11679669



```
QY 136 GYGSYNFSP--IN YKTKITNEQKEFVNNSQAWYQEHGKEEYNGKSFNHTVHNINAN--- 190
Db 257 --GTLNQSQWVITYDTKITT-KQKSYTN-----KATLSGDNLDVAVSRNATVNDYG 303
QY 191 AGIEGT-----VKGLKVLKQDK 208
Db 304 SGGQGTGTPPTPPPVKEPPFPADK 329

RESULT 10
A81888
probable surface fibril protein NMA1200 [imported] - Neisseria meningitidis (strain Z249)
C:Species: Neisseria meningitidis
C>Date: 05-May-2000 #sequence_revision 05-May-2000 #text_change 02-Feb-2001
C:Accession: A81888
R:Parkhill, J.; Achtman, M.; James, K.D.; Bentley, S.D.; Churcher, C.; Klee, S.R.; Morel
; Holroyd, S.; Jagels, K.; Leather, S.; Moule, S.; Mungall, K.; Quail, M.A.; Rajadream,
Nature 404, 502-506, 2000
A:Title: Complete DNA sequence of a serogroup A strain of Neisseria meningitidis Z2491.
A:Reference number: A81775; MUID:20222556; PMID:10761919
A:Accession: A81888
A>Status: preliminary
A:Molecule type: DNA
A:Residues: 1-592 <PAR>
A:Cross-references: GB:ALJ62755; GB:ALJ57959; NID:g7379742; PIDN:CAB84461.1; PID:g737989
A:Experimental source: serogroup A, strain Z2491
A:Genetics:
A:Gene: NMA1200

Query Match 8.9%; Score 99.5; DB 2; Length 592;
Best Local Similarity 21.6%; Pred. No. 11;
Matches 43; Conservative 29; Mismatches 90; Indels 37; Gaps 6;

QY 11 GSDDKVATITSGNKSTN-----VTVHKSAGTSSVFYKTKGDMLPEDTTHVRWF 59
Db 147 GANGKKVNIISDTKGLNFAKETAGTNGDTTVHLNGIGST-----LTDTLAGSSASHV--- 198
QY 60 LNNNEKSVYKDIITKDIQGGQQLDLSTLNINVTGTHSNYSGSAITDEKAFPGSK 119
Db 199 -DAGNQSTHYTRAASIKVLNAGW-----NKGVKGTSTTGQSENVFVRTYDVE 248
QY 120 ITVDNTKNTIDVTIPQGYGSYNSFSINYKTKITNEQKEFVNNSQAWYQEHGKEEYNGKS 179
Db 249 FLSDTKTITTVNVEKDKGKREVKIGAKTSVKEKGKLVTKG-----GKGE-NGSS 300
QY 180 FNHTVHNINANAGIEGTVK 198
Db 301 TDEGEGLVTAKEVIDAVNK 319

RESULT 11
T31105
hypothetical protein 2 - Haemophilus ducreyi
C:Species: Haemophilus ducreyi
C>Date: 22-Oct-1999 #sequence_revision 22-Oct-1999 #text_change 22-Oct-1999
C:Accession: T31105
R:Ward, C.K.; Lumbley, S.R.; Latimer, J.L.; Cope, L.D.; Hansen, E.J.
J. Bacteriol. 180, 6013-6022, 1998
A:Title: Haemophilus ducreyi secretes a filamentous hemagglutinin-like protein.
A:Reference number: Z20984; MUID:99030326; PMID:9811662
A:Accession: T31105
A>Status: preliminary; translated from GB/EMBL/DBJ
A:Molecule type: DNA
A:Residues: 1-4919 <WAR>
A:Cross-references: EMBL:AF057696; NID:g3929021; PID:g3929023; PIDN:AAC79761.1
C:Genetics:
A:Gene: lepA2

Query Match 8.8%; Score 98.5; DB 2; Length 4919;
Best Local Similarity 24.8%; Pred. No. 1.9e+02;
Matches 53; Conservative 34; Mismatches 70; Indels 57; Gaps 12;

QY 15 KVATITSGNKSTNVTVHKSAGTSSVFYKTKGDMLPEDTTHVRWFLNINNEKSVYVSKDIT 74
```

```
Db 1104 RIGKIYAGR---DLTFNKSNAAGKS-----EINRGST-----INVKNKLSYDS-DVS 1146
QY 75 IKDIOGGQQLDLST-----LNINVTGTHSNYSGSAITDEKAFPGSKITVDN 124
Db 1147 FENNQ-SQKVDLYTKIPEAKSDIELTFTKNTGTHPVVLNPKS--NNNEKKYRNS-----EN 1199
QY 125 TKN-----TIDVTIPQGYGSYNS-FSINYKTKITNEQKEFVNNSQAWYQEHGKE 173
Db 1200 TKNFKSGDLINEALSDSAPEAIEAYSGSSNINPVSYLAALGNANNS----- 1250
QY 174 EVNGKSFNHTVHNINANAGIEGTVKGE-LKVLKQ 206
Db 1251 --NPHYLNTALKHILGNGWQDDLLKKQENIKVLKQ 1282

RESULT 12
S33851
fibronectin-binding protein precursor - Streptococcus dysgalactiae
C:Species: Streptococcus dysgalactiae
C>Date: 31-Dec-1993 #sequence_revision 02-Aug-1994 #text_change 15-Oct-1999
C:Accession: S33851; S32631
R:Lindgren, P.E.; McGavin, M.J.; Signaes, C.; Guss, B.; Gurusiddappa, S.; Hoese, M.; Li
Eur. J. Biochem. 214, 819-827, 1993
A:Title: Two different genes coding for fibronectin-binding proteins from Streptococcus
A:Reference number: S33850; MUID:93307299; PMID:8319691
A:Accession: S33851
A:Molecule type: DNA
A:Residues: 1-1117 <LIN>
A:Cross-references: EMBL:Z22151; NID:G288970; PIDN:CAA80122.1; PID:G581643
A:Note: the authors translated the initiation codon TTG for residue 1 as Leu
C:Genetics:
A:Gene: fnbB
A:Start codon: TTG
C:Keywords: fibronectin binding
F:1-27/Domain: signal sequence #status predicted <SIG>
F:28-1117/Product: fibronectin-binding protein #status predicted <MAT>

Query Match 8.7%; Score 98; DB 2; Length 1117;
Best Local Similarity 23.3%; Pred. No. 32;
Matches 51; Conservative 35; Mismatches 85; Indels 48; Gaps 11;

QY 27 NVTVHKSAGTSSVFYKTKGDMLPEDTTH--VRWFLNINNEKSVYVSKDITIK---DOIQ 80
Db 157 NINNVAGSRDTSV-YAKSGVLDLDFDSTGRIVNFWYFNAALNPI-KDEDVEFMTDNL 214
QY 81 GGQQLDLSTLN-----INVVTGTH-SNYSGSAITDEKA-----FPGSKITVDNTK 126
Db 215 QTMTWDIEAVKKDPVWQIRGGYLGTVFSKNGGWIDIEEATKLGIEIIFSGQKLT----- 269
QY 127 NTIDVTIPQGYGSYN-----SFSINYKTKIT---NEQKEFVNNSQAWYQEH 170
Db 270 ----IKPK-WTNYNGLIPYVKPLNQTSSIVKLTAKEKTRMDQTIYVENDSIPKTEG 324
QY 171 GKEEVNGKSFNHTVHNINANAGIEGTVKGLKVLKQDKD 209
Db 325 IEWLDKNQKQFCAKVRQIRSGSWILQVRPDLKVLKFERD 363

RESULT 13
C89874
autolysin [imported] - Staphylococcus aureus (strain N315)
C:Species: Staphylococcus aureus
C>Date: 10-May-2001 #sequence_revision 10-May-2001 #text_change 22-Oct-2001
C:Accession: C89874
R:Kuroda, M.; Ohta, T.; Uchiyama, I.; Baba, T.; Yuzawa, H.; Kobayashi, I.; Cui, L.; Oguc
ma, A.; Mizutani-Ui, Y.; Kobayashi, N.; Sawano, T.; Inoue, R.; Kaito, C.; Sekimizu, K.;
C.; Shiba, T.; Hattori, M.; Ogasawara, N.; Hayashi, H.; Hiramatsu, K.
Lancet 357, 1225-1240, 2001
A:Title: Whole genome sequencing of methicillin-resistant Staphylococcus aureus.
A:Reference number: A89758; MUID:21311952; PMID:11418146
A:Accession: C89874
A>Status: preliminary
```

A;Molecule type: DNA  
A;Residues: 1-1248 <KUR>  
A;Cross-references: GB:BA000018; PID:g13700854; PIDN:BAB42150.1; GSPDB:GNO0149  
A;Experimental source: strain N315  
C;Genetics:  
A;Gene: atl

Query Match 8.7%; Score 98; DB 2; Length 1248;  
Best Local Similarity 21.6%; Pred. No. 37;  
Matches 61; Conservative 39; Mismatches 90; Indels 92; Gaps 16;

Qy 15 KVATITSGNKSTNTVHKSEAGTSSVP-----YKTCG-----MLPEDTH---VRWF 59  
Db 782 KIAQVKNPTGIRASVYKTKAKGAYADRTFVTKERAHGNETYVLLANTSHNPLGWF 841  
Qy 60 ----LNINN--EKSYVSKDITIKDQIQGGQQDLSTLN-INVGTGSHNYSGQSAITDPE 112  
Db 842 NVKDLNVQLGKREKTKTKYTNKSNGLSWPWGTFKNQVILTGN-----IAQGTNATK 897  
Qy 113 KAPPGSKI-----TVDNPK---NTIDVTIP-----OGYGSY---N 141  
Db 898 QVSVGKDVLYGTINRTGWNAKDLTAPTAVKPTTSAAKDYNNTYVVKNGNGYVVTN 957  
Qy 142 SPSINVKTKITNEQ-----KEFVNSQAWYQEHGKEVNGK-----SPNHT 183  
Db 958 SDTAKYSLKAFNEQPPFAVYKEQVINGQTWY--YGKLS-NGKLAWIKSTDLAKELIKYNT 1014  
Qy 184 VHNINAGIEGTVGKELKV-----LKQDKDXTK 211  
Db 1015 GMTLNQVAIQAGLQYKQVQVPGKWTDFANFNDVXHAMDTK 1056

RESULT 14  
C70203  
DNA topoisomerase I (topA) homolog - Lyme disease spirochete  
C;Species: Borrelia burgdorferi (Lyme disease spirochete)  
C;Date: 13-Feb-1998 #sequence\_revision 13-Feb-1998 #text\_change 16-Jul-1999  
C;Accession: C70203  
R;Fraser, C.M.; Casjens, S.; Huang, W.M.; Sutton, G.G.; Clayton, R.; White  
son, D.; Peterson, J.; Kerlavage, A.R.; Quackenbush, J.; Salzberg, S.; Hanson, M.; Vugt,  
; Bowman, C.; Garland, S.; Fujii, C.; Cotton, M.D.; Horst, K.; Roberts, K.; Hatch, B.  
Nature 390, 580-586, 1997  
A;Authors: Smith, H.O.; Venter, J.C.  
A;Title: Genomic sequence of a Lyme disease spirochaete, Borrelia burgdorferi.  
A;Reference number: A70100; MUID:98065943; PMID:9403685  
A;Accession: C70203  
A;Status: preliminary; nucleic acid sequence not shown; translation not shown  
A;Molecule type: DNA  
A;Residues: 1-848 <MLE>  
A;Cross-references: GB:AE001180; GB:AE000783; NID:g2688755; PIDN:AAC67161.1; PID:g268875  
A;Experimental source: strain B31  
C;Superfamily: bacterial type I DNA topoisomerase

Query Match 8.6%; Score 96.5; DB 2; Length 848;  
Best Local Similarity 22.8%; Pred. No. 29;  
Matches 59; Conservative 41; Mismatches 84; Indels 75; Gaps 14;

Qy 19 ITSNGKSTNTVHKSEAGTSSVPFY-----KTGDMLPEDTHVRWFLNINNEKSY 68  
Db 457 IKKGDTFSIVKMKTSBHEKTPAPRYTEASLVQMEKEGGRPSTYSTI---ISTLLREY 513  
Qy 69 VSK-DITIKDQIQGGQDL-----STLNINVTGTH-----SNYYS 103  
Db 514 AFLKNTLPTIKGAVINLLEKFFVLIELNFTSNWEEKLDKIAIGKLDKIKYLSKFYN 573  
Qy 104 GQSAITD-----FEKAFPGSKITVDNTKNTIDVTIPQGYGSY-----NSFS 144  
Db 574 GKKGKLDVTNQLBPKIDSSBFRVIESQKI---ENKNSINYTNIGKYGPYLIFKGNYS 630  
Qy 145 INVTKKITNEOQK-----FVNSQAWYQEHGKEVNGKS---FNHTVH-NI-----NANAG 192  
Db 631 INAKTPLENLYKKDEIKINEKELPNILGVDPLTGLNVIFKNTIYGNIVQLGEDTHAP 690

Qy 193 IEGTVKG---ELKVLKQDK 208  
Db 691 QEYTKKGPKKLKIKAKK 709

RESULT 15  
A53580  
neurexin III beta precursor (clone pB794-7) - bovine  
C;Species: Bos primigenius taurus (cattle)  
C;Date: 12-Apr-1995 #sequence\_revision 12-Apr-1995 #text\_change 09-Dec-2002  
C;Accession: A53580  
R;Ushkaryov, Y.A.; Hata, Y.; Ichtchenko, K.; Moonaw, C.; Afendis, S.; Slaughter, C.A.; S  
J. Biol. Chem. 269, 11987-11992, 1994  
A;Title: Conserved domain structure of beta-neurexins. Unusual cleaved signal sequences  
A;Reference number: A53580; MUID:94216308; PMID:8163501  
A;Accession: A53580  
A;Status: preliminary  
A;Molecule type: mRNA  
A;Residues: 1-392 <USH>  
A;Cross-references: GB:L27864  
C;Superfamily: neurexin; EGF homology  
C;Keywords: alternative splicing

Query Match 8.6%; Score 96; DB 2; Length 392;  
Best Local Similarity 19.6%; Pred. No. 12;  
Matches 47; Conservative 35; Mismatches 74; Indels 84; Gaps 10;

Qy 5 HHHHHGSDDKVATITSGNKSTNTVHK-----EAGTSSVFYVKTGDMLPEDTHVR 57  
Db 52 HEHHPGSK-----HHSVPISYRSPVSLRGHGAGATYFGKSGGLIL-----YT 96  
Qy 58 WFLN-----INNEKSYVSKDITIKDQI-----QG--GOQLDLSTLN 92  
Db 97 WPANDRPSTRDLAVGFSTTVKDGILVRIDSAPGLGDFLQLHIEQKGIGVWFNIGTVDI 156  
Qy 93 NVTGTHSNYSGQSAITDPEKAFPGSKITVDNTKNTIDVTIPQGYGSYNSFSIN--YKTK 150  
Db 157 SIKEERTPVNDGKYHVVRFTNRGNATLQVDN-----WPNVNEHYPTG 198  
Qy 151 ITNEQKKEFVNS-----QAWYQEHGKEVNGKSFNHTVHNINANAGIEGTVKGL 201  
Db 199 NYDNERFQWVKQKIPFKYNRPVEEWLQEKGRQ-----LTIFNTQAIAGGKDKGRL 250

Search completed: July 27, 2004, 10:09:59  
Job time : 17 secs

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GenCore version 5.1.6  
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OM protein - protein search, using sw model

Run on: July 27, 2004, 10:04:21 ; Search time 13 Seconds  
(without alignments)  
845.139 Million cell updates/sec

Title: US-09-813-820-4  
Perfect score: 1122  
Sequence: 1 MRGSHHHHHGSDKVATIT.....GIEGTVKGLKVLKQDKDTK 211

Scoring table: BLOSUM62  
Gapop 10.0 , Gapext 0.5

Searched: 141681 seqs, 52070155 residues

Total number of hits satisfying chosen parameters: 141681

Minimum DB seq length: 0  
Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%  
Maximum Match 100%  
Listing first 45 summaries

Database : SwissProt\_42.\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	ID	Description
1	1048	93.4	1183	1 CNA_STAAU	Q53654 staphylococ
2	106	9.4	1076	1 HSEF_CAVPO	P70106 cavia porce
3	98	8.7	1256	1 ATL_STAAU	P52081 staphylococ
4	96.5	8.6	848	1 TOPI_BORBU	O51768 borrelia bu
5	96	8.6	456	1 NX3B_BOVIN	Q28143 bos taurus
6	96	8.6	704	1 MSN2_YEAST	P33748 saccharomyc
7	95	8.5	666	1 MUR2_ENTHR	P39046 enterococcu
8	95	8.5	2003	1 YDBA_ECOLI	P33666 escherichia
9	93.5	8.3	1296	1 BXB_CLOBO	Q60393 clostridium
10	92.5	8.2	711	1 TOPI_MYCPN	P78032 mycoplasma
11	91.5	8.2	238	1 OPAH_NEIGO	Q04884 neisseria g
12	91	8.1	463	1 GUN_BACSP	P29019 bacillus gp
13	91	8.1	578	1 LIP2_MYCPU	Q50274 mycoplasma
14	90.5	8.1	1169	1 CIGB_BACTZ	Q92426 bacillus th
15	90	8.0	625	1 BSL1_TRIVA	Q8mt12 trichomonas
16	89.5	8.0	416	1 ELT1_CAEEL	P28515 caenorhabdi
17	89	7.9	599	1 LAC2_THACU	Q02075 thanatophor
18	89	7.9	858	1 CYAG_DICDI	Q03101 dictyosteli
19	88.5	7.9	520	1 AMY_EACME	P20845 bacillus me
20	88.5	7.9	576	1 DEAF_DROME	Q24180 drosophila
21	88.5	7.9	717	1 UBPE_YEAST	P36026 saccharomyc
22	87.5	7.8	382	1 Y077_METJA	Q60384 methanococc
23	87.5	7.8	523	1 RPN3_YEAST	P40016 saccharomyc
24	87.5	7.8	626	1 PC11_YEAST	P39081 saccharomyc
25	87.5	7.8	1435	1 DPO3_MYCPU	P47729 mycoplasma
26	87.5	7.8	2867	1 RBP2_PLAUV	Q00799 plasmodium
27	87	7.8	460	1 MURC_THETN	Q8r749 thermonaer
28	87	7.8	668	1 BPBC_BACSU	P42971 bacillus su
29	87	7.8	762	1 SLAP_ACEKI	P22258 acetogenium
30	87	7.8	1310	1 YB35_SCHPO	O14340 schizosacch
31	87	7.8	1433	1 SUBF_BACSU	P16397 bacillus su
32	86.5	7.7	475	1 Y123_MYCPN	P75513 mycoplasma
33	86.5	7.7	678	1 YNC7_YEAST	P53968 saccharomyc

34	86.5	7.7	1694	1 IGA0_HAEIN	P44969 haemophilus
35	86.5	7.7	1702	1 IGA2_HAEIN	P45384 haemophilus
36	86	7.7	437	1 ACMA_LACLC	Q48603 lactococcus
37	86	7.7	617	1 Y237_BUCAL	P57331 buchnera ap
38	86	7.7	986	1 GUNZ_CLOSR	P23659 clostridium
39	85.5	7.6	664	1 AROB_FUSNN	Q8r147 fusobacteri
40	85.5	7.6	850	1 PRSN_PIEBR	Q9g336 pieris bras
41	85.5	7.6	938	1 PM15_CHLPN	Q92883 chlamydia p
42	85.5	7.6	1146	1 AS10_YEAST	P48361 saccharomyc
43	85.5	7.6	1277	1 IF3X_YEAST	Q03690 saccharomyc
44	85	7.6	691	1 PP21_YEAST	P26570 saccharomyc
45	85	7.6	1088	1 VATA_CANTR	P38078 candida tro

ALIGNMENTS

RESULT 1

ID	CNA_STAAU	STANDARD;	PRT; 1183 AA.
AC	Q53654;		
DT	15-DEC-1998 (Rel. 37, Created)		
DT	15-DEC-1998 (Rel. 37, Last sequence update)		
DT	10-OCT-2003 (Rel. 42, Last annotation update)		
DE	Collagen adhesin precursor.		
GN	CNA.		
OS	Staphylococcus aureus.		
OC	Bacteria; Firmicutes; Bacillales; Staphylococcus.		
OX	NCBI_TaxID=1280;		
RN	[1]		
RP	SEQUENCE FROM N.A.		
RC	STRAIN=FDA 574;		
RX	MEDLINE=92165839; PubMed=1311320;		
RA	Patti J.M., Jonsson H., Guss B., Switalski L.M., Wiberg K.,		
RA	Lindberg M., Hoeoek M.;		
RT	"Molecular characterization and expression of a gene encoding a		
RT	Staphylococcus aureus collagen adhesin."		
RL	J. Biol. Chem. 267:4766-4772(1992).		
RN	[2]		
RP	ERRATUM		
RA	Patti J.M., Jonsson H., Guss B., Switalski L.M., Wiberg K.,		
RA	Lindberg M., Hoeoek M.;		
RN	[3]		
RP	COLLAGEN-BINDING DOMAIN.		
RC	STRAIN=FDA 574;		
RX	MEDLINE=94032261; PubMed=8218209;		
RA	Patti J.M., Boles J.O., Hoeoek M.;		
RT	"Identification and biochemical characterization of the ligand		
RT	binding domain of the collagen adhesin from Staphylococcus aureus."		
RL	Biochemistry 32:11428-11435(1993).		
RN	[4]		
RP	X-RAY CRYSTALLOGRAPHY (2.0 ANGSTROMS) OF 169-318.		
RX	MEDLINE=97475225; PubMed=9334749;		
RA	Symersky J., Patti J.M., Carson M., House-Pompeo K., Teale M.,		
RA	Moore D., Jin L., Schneider A., Delucas L.J., Hoeoek M.,		
RA	Narayana S.V.L.;		
RT	"Structure of the collagen-binding domain from a Staphylococcus		
RT	aureus adhesin."		
RL	Nat. Struct. Biol. 4:833-838(1997).		
CC	-!- FUNCTION: MEDIATES ATTACHMENT OF STAPHYOCOCCAL CELLS TO		
CC	COLLAGEN-CONTAINING SUBSTRATA.		
CC	-!- SUBCELLULAR LOCATION: Attached to the cell wall peptidoglycan by		
CC	an amide bond (Potential).		
CC	-----		
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CC	or send an email to license@isb-sib.ch).		
CC	-----		

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DR EMBL; M81736; AAA20874.1; -.
DR PDB; 1AMX; 24-JUN-98.
DR PDB; 1D2O; 27-SEP-00.
DR PDB; 1D2P; 27-SEP-00.
DR InterPro; IPR008966; Adhes_bact.
DR InterPro; IPR008454; Cna_B.
DR InterPro; IPR008970; Cna_B_unit.
DR InterPro; IPR008456; Collagen_bind.
DR InterPro; IPR001899; Gram_pos_anchor.
DR Pfam; PF05738; Cna_B; 7.
DR Pfam; PF05737; Collagen_bind; 1.
DR TIGRFAMs; TIGR01167; LPXTG_anchor; 1.
DR PROSITE; PS50847; GRAM_POS_ANCHORING; FALSE_NEG.
KW Cell wall; Peptidoglycan-anchor; Repeat; Signal; 3D-structure.
FT SIGNAL 1 29 POTENTIAL.
FT CHAIN 30 1154 COLLAGEN ADHESIN.
FT PROPEP 1155 1183 REMOVED BY SORTASE (POTENTIAL).
FT DOMAIN 151 318 COLLAGEN-BINDING.
FT DOMAIN 533 1093 3 X 187 AA APPROXIMATE TANDEM REPEATS.
FT DOMAIN 1093 1157 LYS/PRO-RICH (CELL WALL-SPANNING).
FT REPEAT 533 719 B1.
FT REPEAT 720 906 B2.
FT REPEAT 907 1093 B3.
FT SITE 1151 1155 LPXTG SORTING SIGNAL (POTENTIAL).
FT MOD_RES 1154 1154 AMIDE-LINKED TO CELL WALL (POTENTIAL).
FT STRAND 174 179
FT TURN 182 183
FT TURN 185 186
FT STRAND 187 194
FT TURN 196 197
FT STRAND 201 201
FT STRAND 205 211
FT STRAND 215 228
FT TURN 229 230
FT STRAND 232 234
FT HELIX 239 246
FT TURN 248 249
FT STRAND 251 255
FT TURN 256 259
FT STRAND 260 265
FT HELIX 267 270
FT TURN 271 272
FT STRAND 273 283
FT STRAND 286 287
FT STRAND 290 299
FT STRAND 301 301
FT TURN 302 303
FT STRAND 307 311
FT STRAND 314 317
FT SEQUENCE 1183 AA; 133066 MW; B6A1CC072E575D76 CRC64;

Query Match 93.4%; Score 1048; DB 1; Length 1183;
Best Local Similarity 100.0%; Pred. No. 5.6e-70;
Matches 200; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 12 SDDKATITSGNKSTNVTYHKSEAGTSSVYFKTGDMLPEDTTHRVFLINNEKSYVSK 71
Db 144 SDDKATITSGNKSTNVTYHKSEAGTSSVYFKTGDMLPEDTTHRVFLINNEKSYVSK 203
QY 72 DITIKDQIOGGQDLSTLINVTGTHSNYSYGSQAITDFEAFPGSKITVDNTKNTIDV 131
Db 204 DITIKDQIOGGQDLSTLINVTGTHSNYSYGSQAITDFEAFPGSKITVDNTKNTIDV 263
QY 132 TIPOQGYGSVNSFSINVKITITNEQKQEFVNSQAWYQEHGKEEVNGKSPNHTVHNANA 191
Db 264 TIPOQGYGSVNSFSINVKITITNEQKQEFVNSQAWYQEHGKEEVNGKSPNHTVHNANA 323
QY 192 GIEGTVKGLKVLKQDKTK 211
Db 324 GIEGTVKGLKVLKQDKTK 343

HSEr_CAVPO STANDARD; PRT; 1076 AA.
ID HSEr_CAVPO
AC P70106;
DT 01-NOV-1997 (Rel. 35, Created)
DT 01-NOV-1997 (Rel. 35, Last sequence update)
DT 28-FEB-2003 (Rel. 41, Last annotation update)
DE Heat-stable enterotoxin receptor precursor (GC-C) (Intestinal
DE guanylate cyclase) (EC 4.6.1.2) (STA receptor) (Guanylyl cyclase C).
GN GUCY2C OR GUC2C.
OS Cavia porcellus (Guinea pig).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Rodentia; Hystricognathi; Caviidae; Cavia.
OX NCBI_taxID=10141;
RN [1]
RP SEQUENCE FROM N.A.
RC TISSUE=Colon;
RA Kruhoffer M., Cetin Y., Kaempf U., Forssmann W.-G.;
RL Submitted (AUG-1996) to the EMBL/GenBank/DBJ databases.
CC -!- FUNCTION: RECEPTOR FOR THE E. COLI HEAT-STABLE ENTEROTOXIN (E. COLI
CC ENTEROTOXIN MARKEDLY STIMULATES THE ACCUMULATION OF CGMP IN
CC MAMMALIAN CELLS EXPRESSING GC-C). ALSO ACTIVATED BY THE ENDOGENOUS
CC PEPTIDE GUANYLYN.
CC -!- CATALYTIC ACTIVITY: GTP = 3',5'-cyclic GMP + diphosphate.
CC -!- SUBCELLULAR LOCATION: Type I membrane protein.
CC -!- SIMILARITY: Belongs to the adenylyl cyclase class-4/guanylyl
CC cyclase family.
CC -!- SIMILARITY: Contains 1 protein kinase-like domain.
CC -----
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CC -----
DR EMBL; Z74734; CAA98989.1; -.
DR HSP; Q02846; 1AWL.
DR InterPro; IPR001828; ANF_receptor.
DR InterPro; IPR001054; G_cyclase.
DR InterPro; IPR000719; Prot_kinase.
DR InterPro; IPR001245; Tyr_kinase.
DR Pfam; PF01094; ANF_receptor; 1.
DR Pfam; PF00211; guanylate_cyc; 1.
DR Pfam; PF00069; pkinase; 1.
DR PRINTS; PR00109; TYRKINASE.
DR ProDom; PD000001; Prot_kinase; 1.
DR SMART; SM00044; CYCC; 1.
DR PROSITE; PS00452; GUANYLATE_CYCLASES_1; 1.
DR PROSITE; PS50125; GUANYLATE_CYCLASES_2; 1.
DR PROSITE; PS50011; PROTEIN_KINASE_DOM; 1.
KW Receptor; Transmembrane; Glycoprotein; Lyase; cGMP biosynthesis;
KW Signal.
FT SIGNAL 1 23 POTENTIAL.
FT CHAIN 24 1076 HEAT-STABLE ENTEROTOXIN RECEPTOR.
FT DOMAIN 24 433 EXTRACELLULAR (POTENTIAL).
FT TRANSMEM 434 457 POTENTIAL.
FT DOMAIN 458 1076 CYTOPLASMIC (POTENTIAL).
FT DOMAIN 492 752 PROTEIN KINASE-LIKE.
FT DOMAIN 827 957 GUANYLATE CYCLASE.
FT CARBOHYD 35 35 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 82 82 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 191 191 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 198 198 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 287 287 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 306 306 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 310 310 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 348 348 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 360 360 N-LINKED (GLCNAC. .) (POTENTIAL).
FT CARBOHYD 405 405 N-LINKED (GLCNAC. .) (POTENTIAL).
FT SEQUENCE 1076 AA; 123119 MW; 9B53F16E05E80EB9 CRC64;

Query Match 9.4%; Score 106; DB 1; Length 1076;
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Best Local Similarity 19.6%; Pred. No. 2.6; Matches 47; Conservative 37; Mismatches 78; Indels 78; Gaps 10;

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QY 24 KSTNVTVHKSEAGTSSVFYKGTGMLPDTTHVRWFLN-INNEKSVSKDITIKDQIQG 82
DB 179 KASNLFPKFSWNTSVF--KNG-----TSEHCFTWIINALEAGVSFVSQVLFKEWLRGN 232
QY 83 QQLD-----LSTLNINVTGTH-----SNYSGQSAITDF 111
DB 233 EELQKILKPNRRSNVIVMCGTPQTMESLKIDWTATEDVILVDFNNYILEENATAP- 291
QY 112 EKAFPSKLTVDNKTIDVTITPOGYSVNSFINSYKTKITNEQKQEFVNNQAW-----166
DB 292 -----DYMKNVLVLTLPNGNSTIN-----TSLSKESLQEFDFALAYLDGIL 333
QY 167 -----YOEHGKEVNGCKSFNHTVHNINANAGI-----EGTVKGLKVLKQDKQTK 211
DB 334 LFGHMLKTFERNG-ENTTAHKFAHAFRNLTFFEGSTGPTVLDSDGIDINTVWLLYSVDTK 392
```

## RESULT 3

```
ATL STAAU STANDARD; PRT; 1256 AA.
AC P52081;
DT 01-OCT-1996 (Rel. 34, Created)
DT 01-OCT-1996 (Rel. 34, Last sequence update)
DT 28-FEB-2003 (Rel. 41, Last annotation update)
DE Bifunctional autolysin precursor [includes: N-acetylmuramoyl-L-alanine
DE amidease (EC 3.5.1.28); Mannosyl-glycoprotein endo-beta-N-
DE acetylglucosaminidase (EC 3.2.1.1.96)].
GN ATL.
OS Staphylococcus aureus.
OC Bacteria; Firmicutes; Bacilliales; Staphylococcus.
OX NCBI_TaxID=1280;
RN [1] SEQUENCE FROM N.A., AND SEQUENCE OF 205-214 AND 776-792.
RC STRAIN=RN450;
RX MEDLINE=95116542; PubMed=7816834;
RA Oshida T., Sugai M., Komatsuzawa H., Hong Y.-M., Suganaka H.,
RA Tomasz A.;
RT "A Staphylococcus aureus autolysin that has an N-acetylmuramoyl-L-
RT alanine amidease domain and an endo-beta-N-acetylglucosaminidase
RT domain: cloning, sequence analysis, and characterization."
RL Proc. Natl. Acad. Sci. U.S.A. 92:285-289(1995).
RN [2]
RP SEQUENCE FROM N.A.
RC STRAIN=NCTC 8325-4;
RA Foster S.J.;
RL Submitted (APR-1995) to the EMBL/GenBank/DBJ databases.
CC -!- FUNCTION: ENDOLYOLYSIS OF THE DI-N-ACETYLCHITOBIOSYL UNIT IN
CC HIGH-MANNOSE GLYCOPROTEINS AND GLYCOPROTEINS CONTAINING THE
CC -[(MAN)5(GLCNAc)2]-ASN STRUCTURE. ONE N-ACETYL-D-GLUCOSAMINE
CC RESIDUE REMAINS ATTACHED TO THE PROTEIN; THE REST OF THE
CC OLIGOSACCHARIDE IS RELEASED INTACT.
CC -!- CATALYTIC ACTIVITY: Hydrolyzes the link between N-acetylmuramoyl
CC residues and L-amino acid residues in certain bacterial cell-wall
CC glycopeptides.
CC -!- CATALYTIC ACTIVITY: Endohydrolysis of the di-N-acetylchitobiosyl
CC unit in high-mannose glycopeptides and glycoproteins containing
CC the -[Man(GlcNAc)2]Aen-structure. One N-acetyl-D-glucosamine
CC residue remains attached to the protein; the rest of the
CC oligosaccharide is released intact.
CC -!- SUBCELLULAR LOCATION: Secreted.
CC -!- PTM: UNDERGOES PROTEOLYTIC PROCESSING TO GENERATE THE TWO
CC EXTRACELLULAR LYIC ENZYMES.
CC -!- SIMILARITY: IN THE N-TERMINAL SECTION; BELONGS TO THE N-
CC ACETYLMURAMOYL-L-ALANINE AMIDASE FAMILY 2.
CC -!- SIMILARITY: IN THE C-TERMINAL SECTION; BELONGS TO FAMILY 73 OF
CC GLYCOSYL HYDROLASES.
CC -----
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CC EMBL; D17366; BAA04185.1; -.
CC EMBL; L41499; AAA99982.1; -.
CC InterPro; IPR002502; Amidase_2.
CC Pfam; PF01510; Amidase_2; 1.
CC Pfam; PF01832; Amidase_4; 1.
CC SMART; SM00644; Ami_2; 1.
DR SMART; SM00047; LY22; 1.
KW Cell wall; Hydrolase; Signal; Multifunctional enzyme; Repeat.
FT SIGNAL 1 29
FT CHAIN 30 1256
FT DOMAIN 159 775
FT DOMAIN 776 1256
FT REPEAT 425 589
FT REPEAT 596 758
FT REPEAT 770 932
SQ SEQUENCE 1256 AA; 137384 MW; 2BB76CAA292FDD20 CRC64;
```

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Query Match 8.7%; Score 98; DB 1; Length 1256;
Best Local Similarity 21.6%; Pred. No. 12;
Matches 61; Conservative 39; Mismatches 90; Indels 92; Gaps 16;
QY 15 KVATITSGNKSINVTVHKSEAGTSSVF---YYKTGD-----MLPEDTTH---VRWF 59
DB 790 KIAQVKPNNTGIRASVIEKTXNGAKYADRTFYVTKERAHGNETVLLNNTSHNPLGWF 849
QY 60 ----LNINN--EKSYVSKDITIKDQIQGQQLDLSTLN-INVTGTHSNYSQSAITDFE 112
DB 850 NVKDLNVQNLGKEVKTQKYTVNKSNNGLSMVPMGCTKQVILTGN-----IAQGTFNATK 905
QY 113 KAPPSKI-----TVDNK---NTIDVTIP-----QGYGSY-----N 141
DB 906 QVSVGKDVLYGTINRTGWNNAKDLTAPTAVKPTTSAKDYNVTYVINGNGYVYVTN 965
QY 142 SPSINYKTKITNEQQ---KEFVNNSOAWYQSHGKEVNGK-----SFNHT 183
DB 966 SDATKYSLKAFNEQFPVAVKEQVINGQITWY--YGLKS-NGKLAWIKSTDLAKELIKYNOT 1022
QY 184 VHNINANAGIEGTVKGELKV-----LKQDKDKT 211
DB 1023 GWTLNQVAQIQAGLOYKQVQVRPGKWDKAFNDVKHAMDTK 1064
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## RESULT 4

```
TOP1 BORBU STANDARD; PRT; 848 AA.
AC O51768;
DT 15-DEC-1998 (Rel. 37, Created)
DT 15-DEC-1998 (Rel. 37, Last sequence update)
DT 28-FEB-2003 (Rel. 41, Last annotation update)
DE DNA topoisomerase I (EC 5.99.1.2) (Omega-protein) (Relaxing enzyme)
DE (Untwisting enzyme) (Swivelase).
GN TOPA OR BB0828.
OS Borrelia burgdorferi (Lyme disease spirochete).
OC Bacteria; Spirochaetes; Spirochaetales; Spirochaetaceae; Borrelia.
OX NCBI_TaxID=139;
RN [1] SEQUENCE FROM N.A.
RC STRAIN=ATCC 35210 / B31;
RX MEDLINE=98065943; PubMed=9403685;
RA Fraser C.M., Casjens S., Huang W.M., Sutton G.G., Clayton R.A.,
RA Lathigra R., White O., Ketchum K.A., Dodson R., Hickey E.K., Gwinn M.,
RA Dougherty B., Tomb J.-F., Fleischmann R.D., Richardson D.,
RA Peterson J., Kerlavage A.R., Quackenbush J., Salzberg S., Hanson M.,
RA van Vugt R., Palmer N., Adams M.D., Gocayne J.D., Weidman J.,
RA Utterback T., Watheyl L., McDonald L., Artiach P., Bowman J.,
RA Garland S., Fujii C., Cotton M.D., Horst K., Roberts K., Hatch B.,
RA Smith H.O., Venter J.C.;
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FT DOMAIN 403 456 CYTOPLASMIC (POTENTIAL).
FT DOMAIN 82 282 LAMININ G-LIKE.
FT DOMAIN 41 45 POLY-SER.
FT DOMAIN 311 314 POLY-THR.
FT DOMAIN 317 320 POLY-THR.
FT DOMAIN 369 332 POLY-ALA.
FT CARBOHYD 181 181 N-LINKED (GLCNAC. . .) (POTENTIAL).
FT CARBOHYD 279 279 N-LINKED (GLCNAC. . .) (POTENTIAL).
FT CARBOHYD 323 323 N-LINKED (GLCNAC. . .) (POTENTIAL).
FT VARSPLIC 198 227 Missing (in isoform 2 and isoform 4).
FT VARSPLIC /FTId=VSP_003533.
FT VARSPLIC ANPTGVRVPGASREVIRESSITGMVGVIAAALCILII
FT LUYAMYKYNRDEGSQVDETNYISNSAQSNGLLKEKPP
FT SSKGGHKQKDKKEYV -> ARSNAARSRAALTWTWR
FT LVTFTPIIFISCVHVS (in isoform 3 and
FT isoform 4).
FT /FTId=VSP_003534.
SQ SEQUENCE 456 AA; 50244 MW; E1B546708A8CF2C CRC64;

Query Match 8.6%; Score 96; DB 1; Length 456;
Best Local Similarity 19.6%; Pred. No. 5;
Matches 47; Conservative 35; Mismatches 74; Indels 84; Gaps 10;

Qy 5 HHHHHGSDKVAITSGNKSNTNVTVHKS-----EAGTSSVFFYKTCGDMLPEDTTHVR 57
Db 52 HEHFHFGSK-----HHSVPISYIRSPVSLRGHAGATYFGSGGLIL-----YT 96
Qy 58 WFLN-----INNEKSVYKDTIKDQI-----QG--GQQLDLSTLNI 92
Db 97 WPANDRPSRDLRAVGFSTTVKDGILVIRIDRAGLGDPLQLHIEQKIGVFNIGTVDI 156
Qy 93 NYVTGSHNTSYSGSAITDFEKAFFPGSKITVDNTKNTIDVTIPQGYSYNSFSIN--YTKT 150
Db 157 SIKERTPTVDGKYHVVRTRNGGNATLQVDN-----WPNVHEHYPTG 198
Qy 151 ITNEQKEFVNNS-----QAWYQEHGKEEVNGKSFNHTVHNINANAGIEGTVKGL 201
Db 199 NTDNERFOMVKQIPKPPNRPVVEWLQERGR-----LTFTNTQQAIGGDKGRL 250

RESULT 6
MSN2_YEAST STANDARD; PRT; 704 AA.
AC P33748;
DT 01-PEB-1994 (Rel. 28, Created)
DT 01-PEB-1994 (Rel. 28, Last sequence update)
DT 10-OCT-2003 (Rel. 42, Last annotation update)
DE Zinc finger protein MSN2 (Multicopy suppressor of SNF1 protein 2).
GN MSN2 OR YMR037C OR YW9532.02C.
OS Saccharomyces cerevisiae (Baker's yeast).
OC Eukaryota; Fungi; Ascomycota; Saccharomycotina; Saccharomycetes;
OC Saccharomycetales; Saccharomycetaceae; Saccharomycetes.
OX NCBI_TaxID=4932;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=S288C;
RX MEDLINE=93309420; PubMed=8321194;
RA Estruch F., Carlson M.
RT "Two homologous zinc finger genes identified by multicopy suppression
in a SNF1 protein kinase mutant of Saccharomyces cerevisiae.";
RL Mol. Cell. Biol. 13:3872-3881(1993).
RN [2]
RP SEQUENCE FROM N.A.
RC STRAIN=S288C / AB972;
RX MEDLINE=97313268; PubMed=9169872;
RA Bowman S., Churcher C.M., Badcock K., Brown D., Chillingworth T.,
RA Connor R., Dedman K., Devlin K., Gentles S., Hamlin N., Hunt S.,
RA Jagels K., Lye G., Moule S., Odell C., Pearson D., Rajandream M.A.,
RA Rice P., Skelton J., Walsh S., Whitehead S., Barrell B.G.;
RT "The nucleotide sequence of Saccharomyces cerevisiae chromosome
XIII.";
RL Nature 387:90-93(1997).
RN [3]

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RP CHARACTERIZATION.
RX MEDLINE=96208509; PubMed=8641288;
RA Martinez-Pastor M.T., Marchler G., Schueller C., Marchler-Bauer A.,
RA Ruis H., Estruch F.;
RT "The Saccharomyces cerevisiae zinc finger proteins Men2p and Men4p
are required for transcriptional induction through the stress
response element (STRE).";
RL EMBO J. 15:2227-2235(1996).
RN [4]
RP NUCLEOCYTOPLASMIC SHUTTLING.
RX MEDLINE=22628141; PubMed=12732613;
RA Jacquet M., Renault G., Lallet S., De Mey J., Goldbeter A.;
RT "Oscillatory nucleocytoplasmic shuttling of the general stress
response transcriptional activators Men2 and Men4 in Saccharomyces
cerevisiae.";
RL J. Cell Biol. 161:497-505(2003).
CC -!- FUNCTION: POSITIVE TRANSCRIPTIONAL FACTOR THAT ACTS AS A COMPONENT
OF THE STRESS RESPONSIVE SYSTEM. RECOGNIZES AND BINDS TO THE
STRESS RESPONSE ELEMENT (STRE) WHICH IS INVOLVED IN THE RESPONSE
TO VARIOUS FORMS OF STRESS (HEAT, OXIDATIVE, OSMOTIC, ETC.).
CC INVOLVED IN THE REGULATION OF THE CTT1, DDR2, HSP12 GENES.
CC -!- SUBCELLULAR LOCATION: Cytoplasmic and nuclear.
CC -!- SIMILARITY: Contains 2 C2H2-type zinc fingers.
CC
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CC
CC EMBL; L08838; AAA34806.1; -.
CC EMBL; 248502; CAA88403.1; -.
CC PIR; S39004; S39004.
CC HSP; P08047; ISP1.
CC GerMOnline; 142706; -.
CC TRANSFAC; T01257; -.
CC SGD; S0004640; MSN2.
CC GO; GO:0005829; C:cytosol; IDA.
CC GO; GO:0005634; C:nucleus; IDA.
CC GO; GO:0006950; P:response to stress; IMP.
CC InterPro; IPR007087; Znf_C2H2.
CC Pfam; PF00096; zf-C2H2; 2.
CC ProDom; PD000003; Znf_C2H2; 1.
CC SMART; SM00355; Znf_C2H2; 2.
CC PROSITE; PS00028; ZINC_FINGER_C2H2_1; 2.
CC PROSITE; PS0157; ZINC_FINGER_C2H2_2; 2.
CC DNA-binding; Nuclear protein; Zinc-finger; Metal-binding; Activator;
KW Transcription regulation; Repeat.
FT DOMAIN 1 109 ASP-RICH (ACIDIC).
FT DOMAIN 260 279 ASP-RICH (ACIDIC).
FT ZN_FING 647 665 C2H2-TYPE 1.
FT ZN_FING 676 698 C2H2-TYPE 2.
FT SEQUENCE 704 AA; 77860 MW; EDF6F07446819DF1 CRC64;
SQ
Query Match 8.6%; Score 96; DB 1; Length 704;
Best Local Similarity 25.4%; Pred. No. 8.5;
Matches 64; Conservative 31; Mismatches 97; Indels 60; Gaps 13;

Qy 12 SDDKVAITSGNKSNTNVTVHKSAGTSSVFFYKTCGDMLPEDTTHVRWFNINNEKSYVSK 71
Db 229 SDTNSVNSISNSNSNTGNLN-----SSVFNLSNIDSMLDYVSSDILLNDDDDDTNLSR 284
Qy 72 ----DITKDOIQG-----GQQLDL-----STLNINVTGSHNTSYSGSAIT 109
Db 285 RRFSDV-ITNQPPSMTNSRNSISHSLDLWNHPKINPSNRNTNLNIT-TNSTSSNASPNT 342
Qy 110 DFEKAPPKSKITVDNTKN---TIDVTIPQGYSYN-----SFSI 145
Db 343 TTMNANADSN1-AGNPKNDATIDNELTQILNEYNNFNNDNLGTSTGKNKSKACSSFDA 401
Qy 146 NYTKKITNEQO-KEFVNNSQ---AWYQEHGKEEVNGKSFNHTVHNINANAG----IEGTVKG 199

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Db 402 NAWTKINPQQLQQQLNRVQHKLQTSNHNSSNNKSFNSDLYSRQRASLPIIDSLSY 461  
QY 200 ELKVLKQDKTK 211  
Db 462 DL-VNKQDEDPK 472

RESULT 7  
MUR2\_ENTHR  
ID MUR2\_ENTHR STANDARD; PRT; 666 AA.  
AC P39046;  
DT 01-FEB-1995 (Rel. 31, Created)  
DT 01-FEB-1995 (Rel. 31, Last sequence update)  
DT 16-OCT-2001 (Rel. 40, Last annotation update)  
DE Muramidase-2 precursor (EC 3.2.1.17) (1,4-beta-N-  
DE acetylmuramoylhydrolase) (Peptidoglycan hydrolase) (Pg-hydrolase-2)  
DE (Lysosyme).  
OS Enterococcus hirae.  
OC Bacteria; Firmicutes; Lactobacillales; Enterococcaceae; Enterococcus.  
OX NCBI\_TaxID=1354;

RN [1]  
RP SEQUENCE FROM N.A., AND SEQUENCE OF 50-73.  
RC STRAIN=ATCC 9790;  
RX MEDLINE=92165737; PubMed=1347040;  
RA Chu C.-P., Kariyama R., Daneo-Moore L., Shockman G.D.;  
RT "Cloning and sequence analysis of the muramidase-2 gene from  
RT Enterococcus hirae";  
RL J. Bacteriol. 174:1619-1625 (1992).  
RN [2]  
RP FUNCTION.  
RC STRAIN=ATCC 9790;  
RX MEDLINE=89327152; PubMed=2753858;  
RA Dollinger D.L., Daneo-Moore L., Shockman G.D.;  
RT "The second peptidoglycan hydrolase of Streptococcus faecium ATCC 9790  
RT covalently binds penicillin";  
RL J. Bacteriol. 171:4355-4361 (1989).  
RN [3]  
RP FUNCTION.  
RC STRAIN=ATCC 9790;  
RA Del Mar Lleo M., Caneparl P., Satta G.;

RT "Thermosensitive cell growth mutants of Enterococcus hirae that  
RT elongate at non-permissive temperature are stimulated to divide by  
RT parental autolytic enzymes";  
RL J. Gen. Microbiol. 139:3099-3117 (1993).

CC -!- FUNCTION: MAY WORK IN CONCERT WITH AND POTENTIATE THE PROCESSIVE  
CC HYDROLYTIC ACTION OF MURAMIDASE-1, WHICH REQUIRES BINDING OF THE  
CC ENZYME TO NONREDUCING ENDS OF GLYCAN CHAINS. HYDROLYSIS IN THE  
CC MIDST OF GLYCAN CHAINS WOULD INCREASE THE NUMBER OF BINDING SITES  
CC FOR MURAMIDASE-1. MAY FUNCTION IN FACILITATING SEPTUM FORMATION  
CC AND CELL SEPARATION. ACTIVE ON M.LUTEUS CELL WALLS AND ON E.HIRAE  
CC CELL WALL FRACTIONS, BUT NOT ACTIVE ON E.HIRAE INTACT CELL WALLS.  
CC CAN COVALENTLY BIND PENICILLIN.

CC -!- CATALYTIC ACTIVITY: Hydrolysis of the 1,4-beta-linkages between N-  
CC acetyl-D-glucosamine and N-acetylmuramic acid in peptidoglycan  
CC heteropolymers of the prokaryotes cell walls.

CC -!- SUBCELLULAR LOCATION: Secreted.  
CC -!- DOMAIN: LysM repeats are thought to be involved in peptidoglycan  
CC binding.

CC -!- SIMILARITY: Belongs to family 73 of glycosyl hydrolases.  
CC -!- SIMILARITY: Contains 6 LysM repeats.

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CC -----  
CC EMBL; M77639; AAA24776.1; -.  
CC PIR; A42296; A42296.  
CC HSSP; P22629; 1SWD.

DR InterPro; IPR002901; Amidase 4.  
DR InterPro; IPR000423; Flag\_FlgJ.  
DR InterPro; IPR002482; LysM.  
DR InterPro; IPR000437; Prok\_lipoprot\_S.  
DR Pfam; PF01832; Amidase\_4; 1.  
DR Pfam; PF01476; LysM; 6.  
DR PRINTS; PR01002; FLGFLGJ.  
DR SMART; SM00257; LysM; 6.  
DR SMART; SM00047; LysM; 1.  
DR SMART; SM00047; LysM; 6.  
KW Hydrolase; Glycosidase; Bacteriolytic enzyme; Cell wall;  
KW Cell division; Septation; Repeat; Signal.

FT SIGNAL 1 49  
FT CHAIN 50 666 MURAMIDASE-2.  
FT REPEAT 257 299 LYSM 1.  
FT REPEAT 338 380 LYSM 2.  
FT REPEAT 414 456 LYSM 3.  
FT REPEAT 489 531 LYSM 4.  
FT REPEAT 565 607 LYSM 5.  
FT REPEAT 623 665 LYSM 6.  
SQ SEQUENCE 666 AA; 70670 MW; FFOA7FAFCD810BA3 CRC64;

Query Match 8.5%; Score 95; DB 1; Length 666;  
Best Local Similarity 24.0%; Pred. No. 9.4;  
Matches 49; Conservative 28; Mismatches 81; Indels 46; Gaps 11;

QY 20 TSGNKSNTVTVHSEAGTSSVYYKTGDMLEPDTH-----VRFLNINNEKSYVS 70  
Db 242 TSGNKGSGSATT-----TGTTYTVKSGDSV-WGISHSGITMAQLIEW-NNIKNNFIYPG 293  
QY 71 KDIITKQIQGGQQLDLSTLNI-----NVTG-THSNYSGQSAITDFEKAFFGSKITV 122  
Db 294 QKLTIK-----GGQAGSSTTNTGNASSGNTSGNTSGTGQAT-----GAKYTV 340  
QY 123 DNTKNTIDVTIPQGVGNSPFSINYKTKITNEQKQFVNNSQAWYQEHGKEVNGKSPNH 182  
Db 341 KSGDSVWKIANDHG-----ISMQLIE-WNNIKNNFYPCQLVWVSKGSSASGTSNT 393  
QY 183 TVHNINANAGIEG-TVKGLKVLK 205  
Db 394 STGNTSSNTANTGTTSGSTYTVK 417

## RESULT 8

YDBA\_ECOLI  
ID YDBA\_ECOLI STANDARD; PRT; 2003 AA.  
AC P33666; P76087; P76088; P76856; P76857; P76859;  
DT 01-FEB-1994 (Rel. 28, Created)  
DT 16-OCT-2001 (Rel. 40, Last sequence update)  
DE Hypothetical protein ydBA.  
GN YDBA OR B1401/B1405.  
OS Escherichia coli.  
OC Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales;  
OC Enterobacteriaceae; Escherichia.  
OX NCBI\_TaxID=562;  
RP [1]  
RP SEQUENCE FROM N.A.  
RC STRAIN=K12 / MG1655;  
RX MEDLINE=97426617; PubMed=9278503;  
RA Blattner F.R., Plunkett G. III, Bloch C.A., Perna N.T., Burland V.,  
RA Riley M., Collado-Vides J., Glasner J.D., Rode C.K., Mayhew G.F.,  
RA Gregor J., Davis N.W., Kirkpatrick H.A., Goeden M.A., Rose D.J.,  
RA Mau B., Shao Y.;  
RA "The complete genome sequence of Escherichia coli K-12.";  
RL Science 277:1453-1474 (1997).  
RN [2]  
RP SEQUENCE FROM N.A.  
RC STRAIN=K12;  
RX MEDLINE=97251357; PubMed=9097039;  
RA Alba H., Baba T., Fujita K., Hayashi K., Inada T., Isono K., Itoh T.,  
RA Kasai H., Kishimoto K., Kimura S., Kitagawa M., Kitagawa M.,  
RA Makino K., Miki T., Mizobuchi K., Mori H., Mori T., Motomura K.,  
RA Nakade S., Nakamura Y., Nashimoto H., Nishio Y., Oshima T., Saito N.,





QY 148 -KTKITNEOQKEF-----VNNQAW-YQHGKE-----EVNGKS----- 179  
 Db 944 NNNDIQYLYNYHTIISIKDNGSKVSIKGNRIITWLDVNAKSIFFEYSIKDNISD 1003  
 QY 180 -----FNHTVHNI-----NANAGIEGTGKGLVKLQDK 208  
 Db 1004 YINKWFSITITDRNLGNANIYINGSLKKSEKILNLD 1040

## RESULT 10

TOPI\_MYCPN  
 ID TOPI\_MYCPN STANDARD; PRT; 711 AA.  
 AC P78032;  
 DT 01-NOV-1997 (Rel. 35, Created)  
 DT 01-NOV-1997 (Rel. 35, Last sequence update)  
 DT 28-FEB-2003 (Rel. 41, Last annotation update)  
 DE DNA topoisomerase I (EC 5.99.1.2) (Omega-protein) (Relaxing enzyme)  
 DE (Untwisting enzyme) (Swivelase).  
 GN TOPA OR MPN261 OR MP572.  
 OS Mycoplasma pneumoniae.  
 OC Bacteria; Firmicutes; Mollicutes; Mycoplasmataceae; Mycoplasma.  
 OX NCBI\_TaxID=2104;  
 RN [1]  
 RP SEQUENCE FROM N.A.  
 RC STRAIN=ATCC 29342 / M129;  
 RX MEDLINE=97105885; PubMed=8948633;  
 RA Himmelreich R., Hilbert H., Plagens H., Pirkel E., Li B.-C.,  
 RA Herrmann R.;  
 RT "Complete sequence analysis of the genome of the bacterium Mycoplasma  
 RT pneumoniae.";  
 RL Nucleic Acids Res. 24:4420-4449(1996).  
 CC -!- FUNCTION: The reaction catalyzed by topoisomerases leads to the  
 CC conversion of one topological isomer of DNA to another.  
 CC -!- CATALYTIC ACTIVITY: ATP-independent breakage of single-stranded  
 CC DNA, followed by passage and rejoining.  
 CC -!- SUBUNIT: Monomer (By similarity).  
 CC -!- MISCELLANEOUS: When a topoisomerase transiently breaks a DNA  
 CC backbone bond, it simultaneously forms a protein-DNA link, in  
 CC which a tyrosyl oxygen in the enzyme is joined to a DNA phosphorus  
 CC at one end of the enzyme-severed DNA strand.  
 CC -!- SIMILARITY: Belongs to the prokaryotic type I/III topoisomerase  
 CC family.  
 CC -----  
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 CC -----  
 DR EMBL; A5000056; AAB96220.1; -.  
 DR F1R; S73898; S73898.  
 DR HSP; P06612; 1ECL.  
 DR InterPro; IPR005733; DNA\_topi\_bact.  
 DR InterPro; IPR000380; DNA\_topoisomerase.  
 DR InterPro; IPR003601; DNA\_topi\_ATP\_bind.  
 DR InterPro; IPR003602; DNATopi\_DNA\_bind.  
 DR InterPro; IPR006171; Toprim\_dom.  
 DR InterPro; IPR006154; Toprim\_sub.  
 DR Pfam; PF01131; Topoisom\_bac; 1.  
 DR Pfam; PF01751; Toprim; 1.  
 DR Pfam; PF01396; zf-C4\_Topoism; 2.  
 DR PRINTS; PR00417; PRFISMRASE1.  
 DR SMART; SM00437; TOPIAC; 1.  
 DR SMART; SM00436; TOPIBC; 1.  
 DR SMART; SM00493; TOPRIM; 1.  
 DR TIGRPFAM; TIGR01051; topA\_bact; 1.  
 DR PROSITE; PS00396; TOPOISOMERASE\_I\_PROK; 1.  
 KW Repeat; Complete proteome.  
 FT ZN\_FING 624 652 C4-TYPE 1.  
 FT ZN\_FING 673 702 C4-TYPE 2.

FT ACT SITE 340 340 DNA CLEAVAGE (BY SIMILARITY).  
 SQ SEQUENCE 711 AA; 81965 MW; 168863588B06582E CRC64;  
 Query Match 8.2%; Score 92.5; DB 1; Length 711;  
 Best Local Similarity 25.4%; Pred. No. 16;  
 Matches 49; Conservative 32; Mismatches 65; Indels 47; Gaps 12;  
 QY 54 THVRWFLNINNE-----KSYV-----SKDITIK-DOIOGGQOOLDLSTLNI 92  
 Db 436 THIR-FINDGNKFYASSKSLVFDGVRKIYEFENKESNDLYIDDKRVGRFMAKDRI 494  
 QY 93 NVTGTH-SNYYSQSAITDFEKAFFGSKITVDNTKNTI-DVTIPQGYGSYN--SFSINYK 148  
 Db 495 TARQTHPAARYTQASLIEALEK----SNIGRPSTVNTWASVNLDRGYASLNKHAHVHTQL 550  
 QY 149 TKITNEQ-----KQEFVNNQAWQVQEGKEEV--NGKSFNHTVHNIANAGIE-GT 196  
 Db 551 GEQVNEELSHFGKIINKFTKN-----MEKSLDEIAENKKNYQFELDRDFWSNFKEVKL 605  
 QY 197 VKGELKVLKQDKD 209  
 Db 606 AEGSIQVRVKEKE 618  
 RESULT 11  
 OPAH\_NEIGO  
 ID OPAH\_NEIGO STANDARD; PRT; 238 AA.  
 AC Q04884;  
 DT 01-OCT-1993 (Rel. 27, Created)  
 DT 01-OCT-1993 (Rel. 27, Last sequence update)  
 DT 15-DEC-1998 (Rel. 37, Last annotation update)  
 DE Opacity protein OPA60 precursor (Fragment).  
 GN OPAH.  
 OS Neisseria gonorrhoeae.  
 OC Bacteria; Proteobacteria; Betaproteobacteria; Neisseriales;  
 OC Neisseriaceae; Neisseria.  
 OC NCBI\_TaxID=485;  
 RN [1]  
 RP SEQUENCE FROM N.A.  
 RC STRAIN=MS11 / F3;  
 RX MEDLINE=93178439; PubMed=8440254;  
 RA Kupsch E.-M., Knepper B., Kuroki T., Heuer I., Meyer T.F.;  
 RT "Variable opacity (Opa) outer membrane proteins account for the cell  
 RT tropisms displayed by Neisseria gonorrhoeae for human leukocytes and  
 RT epithelial cells";  
 RL EMO J. 12:641-650(1993).  
 RN [2]  
 RP SEQUENCE FROM N.A.  
 RC STRAIN=MS11 / V18;  
 RX MEDLINE=92114767; PubMed=1815562;  
 RA Bhat K.S., Gibbs C.P., Barrera O., Morrison S.G., Jaehnig F.,  
 RA Stern A., Kupsch E.-M., Meyer T.F., Swanson J.;  
 RT "The opacity proteins of Neisseria gonorrhoeae strain MS11 are  
 RT encoded by a family of 11 complete genes.";  
 RL Mol. Microbiol. 5:1889-1901(1991).  
 RN [3]  
 RP ERRATUM  
 RX MEDLINE=92261323; PubMed=1584024;  
 RA Bhat K.S., Gibbs C.P., Barrera O., Morrison S.G., Jaehnig F.,  
 RA Stern S., Kupsch E.-M., Meyer T.F., Swanson J.;  
 RL Mol. Microbiol. 6:1073-1076(1992).  
 CC -!- FUNCTION: IMPLICATED IN A NUMBER OF ADHERENCE FUNCTIONS. OPA  
 CC PROTEINS ARE IMPLICATED IN PATHOGENESIS AND ARE SUBJECT TO PHASE  
 CC VARIATION.  
 CC -!- SUBCELLULAR LOCATION: Outer membrane.  
 CC -----  
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```
CC -----
DR EMBL; Z18939; CAA79372.1; -.
DR EMBL; X60711; CAA43121.1; -.
DR PIR; S16619; S16619.
DR InterPro; IPR003394; Porin_opacity.
DR Pfam; PF02462; Opacity; 1. family; Signal.
KW Outer membrane; Multigene family; Signal.
FT NON_TER 1
FT SIGNAL <1 1 POTENTIAL.
FT CHAIN <1 1 OPACITY PROTEIN OPA60.
FT VARIAT 2 >238 SED -> MLKA (IN MS11 / V18).
FT VARIAT 234 234 V -> M (IN MS11 / V18).
FT NON_TER 238 238
FT SEQUENCE 238 AA; 27073 MW; 883A3560C2DF1B9F CRC64;
SQ
Query Match 8.2%; Score 91.5; DB 1; Length 238;
Best Local Similarity 24.7%; Pred. No. 4.7;
Matches 45; Conservative 35; Mismatches 69; Indels 33; Gaps 9;
QY 4 SHHHHSGDDKVATITSGNKTNTVHKSEAGTSSVF-----YKTKGMLPEDTTHVR 57
DB 16 AVAYEHITHDYPTAPNKKKISTVSDYFRNIRTSVHPRVSVGVDFGCGWRIAADYARVR 75
QY 58 -WFLNNNEKSYVSKDITIKDQIQGGQQLDLSTLNI--NVTKTHSNY--YSGQSA- 112
DB 76 KW-----NNNKYSYNIENVRIRKENGIRIDRKTENQENGTFHVVSLGLSAYIDFQINDK 130
QY 113 -KAPPGSKT-----TVNTKNTIDV-TIPQG--YGSYNSFSGINVKTKITNEQQKEFVN 161
DB 131 FRPYIGARVAYGHVRHSIDSKTKTIEVTIVPSNAPNGAVTTNTDPT-----QNDYOS 184
QY 162 NS 163
DB 185 NS 186
RESULT 12
GUN_BACSP STANDARD; PRT; 463 AA.
AC P29019;
DT 01-DEC-1992 (Rel. 24, Created)
DT 01-DEC-1992 (Rel. 24, Last sequence update)
DT 28-FEB-2003 (Rel. 41, Last annotation update)
DE Endoglucanase precursor (EC 3.2.1.4) (Endo-1,4-beta-glucanase)
DE (Cellulase) (Endo-K).
OS Bacillus sp. (strain KSM-330).
OC Bacteria; Firmicutes; Bacillales; Bacillaceae; Bacillus.
OX NCBI_TaxID=1409;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=92121880; PubMed=1770347;
RA Ozaki K., Sumitomo N., Ito S.;
RT "Molecular cloning and nucleotide sequence of the gene encoding an
RT endo-1,4-beta-glucanase from Bacillus sp. KSM-330.";
RL J. Gen. Microbiol. 137:2299-2305 (1991).
RN [2]
RP SEQUENCE OF 56-75, AND CHARACTERIZATION.
RX MEDLINE=91259037; PubMed=2045781;
RA Ozaki K., Ito S.;
RT "Purification and properties of an acid endo-1,4-beta-glucanase from
RT Bacillus sp. KSM-330.";
RL J. Gen. Microbiol. 137:41-48 (1991).
CC -1- FUNCTION: THIS ACID ENDOGLUCANASE IS ACTIVE OVER AN EXTREMELY
CC NARROW RANGE OF PH VALUES, BETWEEN 4.5 AND 6.5, WITH AN OPTIMUM
CC PH AT 5.2.
CC -1- CATALYTIC ACTIVITY: Endohydrolysis of 1,4-beta-D-glucosidic
CC linkages in cellulose, lichenin and cereal beta-D-glucans.
CC -1- PTM: THE N- AND THE C-TERMINUS MAY BE SUBJECTED TO PROTEOLYSIS.
CC -1- MISCELLANEOUS: ONE TRP RESIDUE HAS BEEN PROVED TO BE INVOLVED IN
CC THE MECHANISM OF ACTION OF ENDO-K.
CC -1- SIMILARITY: Belongs to cellulase family D (family 8 of glycosyl
CC hydrolases).
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CC -----
DR EMBL; M68872; AAA22409.1; -.
DR PIR; A44808; A44808.
DR InterPro; IPR002037; Glyco_hydro_8.
DR InterPro; IPR008928; Glyco_trans_6hp.
DR Pfam; PF01270; Glyco_hydro_8; 1.
DR PRINTS; PR00735; GLHYDRLASE8.
DR PROSITE; PS00812; GLYCOSYL HYDROL F8; 1.
KW Cellulose degradation; Hydrolase; Glycosidase; Signal.
FT SIGNAL 1 27 POTENTIAL.
FT PROPEP 28 55 POTENTIAL.
FT CHAIN 56 463 ENDOGLUCANASE.
FT ACT_SITE 130 130 PROTON DONOR (BY SIMILARITY).
FT ACT_SITE 191 191 NUCLEOPHILE (POTENTIAL).
FT SEQUENCE 463 AA; 51882 MW; 407FA54F5236C59E CRC64;
SQ
Query Match 8.1%; Score 91; DB 1; Length 463;
Best Local Similarity 21.2%; Pred. No. 12;
Matches 53; Conservative 31; Mismatches 64; Indels 102; Gaps 13;
QY 7 HHHHSGDDKV-----ATITSGNKTNTV---VHKSEAGTSSVYKTKGMLPED--T 53
DB 204 HKWGSSGKINYLKSAQNMTGKIKASNVTKNGNLGWDGDKSTF-----DTRSDWNM 258
QY 54 THVRWFLNNNEKSYVSKDITIKDQIQGGQQLDLSTLNI--NVTKTHSNY--YSGQSA- 107
DB 259 SHLRAFYEFTGDKTW-----LNVIDNLYNTYNTNFTNKYSPKTKGL 297
QY 108 ITDP--EKAFFGSKITVDNTKNT-----IDVTIPOGVGSYN-----SPS 144
DB 298 ISDFVYVKNPPQAPKDFLDESKYDTSYYYNASRVPLRVIMDYAMYGEKRGKVISDKVATM 357
QY 145 INYKTK-----ITNEQOKFEVNNQAWYQEHG 171
DB 358 IKSCTKGNPSKIVDGYKLDGTNIGDYPYAVYVSPFIAAGTTNSKNQEWNSGWDW-MKNK 416
QY 172 KEEVNGKSPN 181
DB 417 KESYFSDSYN 426
RESULT 13
LIPA_MYCPU STANDARD; PRT; 578 AA.
ID LIPA_MYCPU
AC Q50274; Q98Q39;
DT 16-OCT-2001 (Rel. 40, Created)
DT 16-OCT-2001 (Rel. 40, Last sequence update)
DT 10-OCT-2003 (Rel. 42, Last annotation update)
DE Lipoprotein A precursor.
GN LIPA OR MYPU 5300.
OS Mycoplasma pulmonis.
OC Bacteria; Firmicutes; Mollicutes; Mycoplasmataceae; Mycoplasma.
OX NCBI_TaxID=2107;
RN [1]
RP SEQUENCE FROM N.A.
RX STRAIN=KD735-15;
RX MEDLINE=20245550; PubMed=10781561;
RA Shen X., Gumulak J., Yu H., French C.T., Zou N., Dybvig K.;
RT "Gene rearrangements in the vsa locus of Mycoplasma pulmonis.";
RL J. Bacteriol. 182:2900-2908 (2000).
RN [2]
RP SEQUENCE FROM N.A.
RX STRAIN=UAB CT1P;
RX MEDLINE=21267165; PubMed=11353084;
RA Chambaud I., Heilig R., Ferris S., Barbe V., Samson D., Galisson F.,
RA Moszer I., Dybvig K., Wroblewski H., Viari A., Rocha E.P.C.,
```

RA Blanchard A.;  
RT "The complete genome sequence of the murine respiratory pathogen  
RT Mycoplasma pulmonis";  
RL Nucleic Acids Res. 29:2145-2153(2001).  
[3]  
RN PARTIAL SEQUENCE FROM N.A.  
RC STRAIN=KD735-15;  
RX MEDLINE=96414471; PubMed=8817492;  
RA Bhugra B., Voelker L.L., Zou N., Yu H., Dybvig K.;  
RT "Mechanism of antigenic variation in Mycoplasma pulmonis: interwoven,  
RT site-specific DNA inversions";  
RL Mol. Microbiol. 18:703-714(1995).  
CC -!- SUBCELLULAR LOCATION: Attached to the membrane by a lipid anchor  
CC (Potential).  
CC -!- SIMILARITY: BELONGS TO THE PULMONIS LIPAB LIPOPROTEIN FAMILY.  
CC  
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CC  
CC EMBL; U23947; AAB41030.2; -.  
DR EMBL; AL445565; CAC13703.1; -.  
DR PIR; B90578; B90578.  
DR MyPulist; NYPU\_5300; -.  
DR InterPro; IPR007326; Lipoprotein\_17.  
DR InterPro; IPR000437; Prok\_lipoprot\_8.  
DR Pfam; PF04200; Lipoprotein\_17; 1.  
DR PROSITE; PS00013; PROKAR LIPOPROTEIN; 1.  
KW Lipoprotein; Membrane; Signal; Complete proteome; Palmitate.  
FT SIGNAL 1 27  
FT CHAIN 28 578 Lipoprotein A.  
FT LIPID 28 28 N-palmitoyl cysteine (Potential).  
FT LIPID 28 28 S-diacylglycerol cysteine (Potential).  
FT DOMAIN 82 165 ASN-RICH.  
FT VARIANT 7 7 K -> R (IN STRAIN KD735-15).  
FT VARIANT 41 41 N -> S (IN STRAIN KD735-15).  
FT VARIANT 110 110 N -> D (IN STRAIN KD735-15).  
FT VARIANT 127 127 S -> A (IN STRAIN KD735-15).  
FT VARIANT 547 547 Y -> N (IN STRAIN KD735-15).  
SQ SEQUENCE 578 AA; 66219 MW; B533ED3467005D89 CRC64;  
  
Query Match 8.1%; Score 91; DB 1; Length 578;  
Best Local Similarity 20.2%; Pred. No. 16;  
Matches 49; Conservative 33; Mismatches 103; Indels 58; Gaps 6;  
  
QY 14 DKVATITSGNK-STNVTVHKSEAGTSSVFPYKTDGMLPDDTHVRWFLNNNEKSYVSKD 72  
DB 61 DKDKRVQDNKSTKAVNSQTSQKTSQNTKDDSKTSNLTNQNSSTNTSK 120  
QY 73 ITIKDIQGGQQLDLSTLNINVTGTHSNYSQSAITDFEAPGSKITVDNTKNTIDVT 132  
DB 121 IQENKQ----SQDKNTSAVNSALE-----KQTKNDENISLVNSKDTNVLKNDKVA 170  
QY 133 IPQGVG---SYNSFSINVKTKITNQCKEFVNSQ---AWYQE----- 169  
DB 171 LAKDSDKEKSKNSNLNLTAPVENQKNNEVDKDKALQWQMKQKESASILESFSYDQTS 230  
QY 170 -----HGKEVNGKSNFHTVHNINAGIEGTGVKGLVK 205  
DB 231 LSLTFKEGMPGLEVVVLKLENLDSHEEKEISFKTNGKVQNVLLTSSNLTSGKWKIKFS 290  
QY 206 QDK 208  
DB 291 FDK 293  
  
RESULT 14  
C1GB\_BACTZ STANDARD; PRT; 1169 AA.  
ID C1GB\_BACTZ

Q9ZAZ6;  
30-MAY-2000 (Rel. 39, Created)  
30-MAY-2000 (Rel. 39, Last sequence update)  
16-OCT-2001 (Rel. 40, Last annotation update)  
DE Pesticidal crystal protein cryIgb (insecticidal delta-endotoxin  
DE CryIgb(b)) (Crystalline entomocidal protoxin) (133 kDa crystal protein).  
GN CRV1GB OR CRV1GB(B) OR CRVH2.  
OS Bacillus thuringiensis (subsp. wuhanensis).  
OC Bacteria; Firmicutes; Bacillales; Bacillaceae; Bacillus.  
OX NCBI\_TaxID=52024;  
[1]  
RP SEQUENCE FROM N.A.  
RC STRAIN=HD-525;  
RX MEDLINE=20153386; PubMed=10688690;  
RA Kuo W.-S., Lin J.-H., Tzeng C.-C., Kao S.-S., Chak K.-F.;  
RT "Cloning of two new cry genes from Bacillus thuringiensis subsp.  
RT wuhanensis strain";  
RL Curr. Microbiol. 40:227-232(2000).  
CC -!- FUNCTION: PROMOTES COLLOIDSMOTIC LYSIS BY BINDING TO THE MIDGUT  
CC EPITHELIAL CELLS OF LEPIDOPTERAN LARVAE. TOXIC TO PIERIS RAPAE.  
CC -!- DEVELOPMENTAL STAGE: The crystal protein is produced during  
CC sporulation and is accumulated both as an inclusion and as part of  
CC the spore coat.  
CC -!- MISCELLANEOUS: Toxic segment of the protein is located in the N-  
CC terminus.  
CC -!- SIMILARITY: Belongs to the delta endotoxin family.  
CC  
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CC  
CC EMBL; U07025; AAD10291.1; -.  
DR HSSP; P02965; 1CIY.  
DR InterPro; IPR001178; Endotoxin.  
DR InterPro; IPR005638; Endotoxin\_C.  
DR InterPro; IPR005639; endotoxin\_N.  
DR InterPro; IPR008979; Gal\_bind\_like.  
DR Pfam; PF00555; endotoxin; 1.  
DR Pfam; PF03944; endotoxin; 1.  
DR Pfam; PF03945; endotoxin\_N; 1.  
DR Toxin; Sporulation.  
KW SEQUENCE 1169 AA; 132904 MW; D1EFC1508A8B10BD CRC64;  
  
Query Match 8.1%; Score 90.5; DB 1; Length 1169;  
Best Local Similarity 24.5%; Pred. No. 41;  
Matches 36; Conservative 17; Mismatches 81; Indels 13; Gaps 5;  
  
QY 28 VTVHKSEAGTSSVFPYKTDGMLPDDTHVRWFLNNNEKSYVSKDITIKDIQGGQQLDL 87  
DB 1021 VTAYKEGVGECVTIHBI-----EDNTDELKFSNCVEEIIYPNNVTVCNDYTATQEEYE- 1074  
QY 88 STLINVTGTHSNYSQSAITDFEAPGSKITVDNTKNTIDVTIPQGYGSYNSFSINY 147  
DB 1075 GTYTSRNRGVDGAVESNSVPADYASAYEEKAYTGDRRDNTCSN--RGYGDYTPLPAGY 1132  
QY 148 KTKITNEQKQEFVNSQAWYQEHGKE 174  
DB 1133 VTK----ELEYFPETDKVWI-EIGETE 1154  
  
RESULT 15  
BSL1\_TRIVA STANDARD; PRT; 625 AA.  
ID BSL1\_TRIVA  
AC Q8MTI2;  
DT 15-MAR-2004 (Rel. 43, Created)  
DT 15-MAR-2004 (Rel. 43, Last sequence update)  
DT 15-MAR-2004 (Rel. 43, Last annotation update)  
DE Putative surface protein bepA-like (TvBspA-like-625).  
GN BSPALL1.



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GenCore version 5.1.6  
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OM protein - protein search, using sw model

Run on: July 27, 2004, 10:05:01 ; Search time 39 Seconds  
(without alignments)  
1707.034 Million cell updates/sec

Title: US-09-813-820-4

Perfect score: 1122

Sequence: 1 MRGSHHHHHGSDKVATIT.....GIEGTVKGLKVLKQDKTK 211

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 1017041 segs, 315518202 residues

Total number of hits satisfying chosen parameters: 1017041

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database :

- 1: sp\_archaea.\*
- 2: sp\_bacteria.\*
- 3: sp\_fungi.\*
- 4: sp\_human.\*
- 5: sp\_invertebrate.\*
- 6: sp\_mammal.\*
- 7: sp\_mhc.\*
- 8: sp\_organelle.\*
- 9: sp\_phage.\*
- 10: sp\_plant.\*
- 11: sp\_rodent.\*
- 12: sp\_virus.\*
- 13: sp\_vertebrate.\*
- 14: sp\_unclassified.\*
- 15: sp\_rvirus.\*
- 16: sp\_bacteriapi.\*
- 17: sp\_archaeap.\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	ID	Description
1	1035	92.2	1183	16 Q8NUH0	Q8nuh0 staphylococ
2	491	43.8	721	2 Q84BK5	Q84bk5 enterococcu
3	463.5	41.3	657	2 Q84OA5	Q84oa5 streptococc
4	462.5	41.2	221	2 Q8KRZ5	Q8krz5 streptococc
5	182.5	16.3	1151	2 Q7WWP3	Q7wvp3 arcanobacte
6	182	16.2	319	2 Q9F859	Q9f859 enterococcu
7	181	16.1	721	2 Q9F866	Q9f866 enterococcu
8	179	16.0	627	2 Q9F863	Q9f863 enterococcu
9	179	16.0	721	2 Q9F867	Q9f867 enterococcu
10	178	15.9	319	2 Q9F870	Q9eu70 enterococcu
11	178	15.9	319	2 Q9F857	Q9f857 enterococcu
12	178	15.9	319	2 Q9F856	Q9f856 enterococcu
13	178	15.9	458	2 Q9XBQ7	Q9xbq7 enterococcu
14	178	15.9	580	2 Q9F862	Q9f862 enterococcu
15	178	15.9	580	2 Q9F861	Q9f861 enterococcu
16	178	15.9	627	2 Q9F864	Q9f864 enterococcu

17	178	15.9	674	2	Q9F865	Q9f865 enterococcu
18	178	15.9	674	16	Q836L2	Q836l2 enterococcu
19	177.5	15.8	319	2	Q9F858	Q9f858 enterococcu
20	176.5	15.7	787	2	Q83VG5	Q83vg5 erysipeloth
21	175	15.6	296	2	Q9F860	Q9f860 enterococcu
22	175	15.6	319	2	Q9EU80	Q9eu80 enterococcu
23	175	15.6	319	2	Q84099	Q84099 streptococ
24	128	11.4	1991	2	Q83VG6	Q83vg6 erysipeloth
25	121	10.8	3381	5	Q812V4	Q812v4 plasmodium
26	119	10.6	336	16	Q9CHP4	Q9chp4 lactococcus
27	118	10.5	1361	5	Q8IBV5	Q8ibv5 plasmodium
28	116.5	10.4	1824	5	Q815A3	Q815a3 plasmodium
29	116	10.3	642	16	Q898I0	Q898i0 clostridium
30	114.5	10.2	1335	5	Q814Y7	Q814y7 plasmodium
31	114.5	10.2	1987	16	Q839R9	Q839r9 enterococcu
32	109.5	9.8	2770	5	Q81LV0	Q81lv0 plasmodium
33	108	9.6	723	16	Q8XPI1	Q8xpi1 clostridium
34	107.5	9.6	1231	16	Q88ZE8	Q88ze8 lactobacill
35	107.5	9.6	2577	5	Q81AMI	Q81aml plasmodium
36	107	9.5	321	16	Q896O5	Q896o5 clostridium
37	107	9.5	358	16	Q931E7	Q931e7 staphylococ
38	105.5	9.4	598	2	Q9JPR7	Q9jpr7 neisseria m
39	105	9.4	3135	5	Q813B5	Q813b5 plasmodium
40	104.5	9.3	1318	5	Q95PH4	Q95ph4 dictyosteli
41	104.5	9.3	2488	5	Q813R5	Q813r5 plasmodium
42	104	9.3	961	5	Q81KT9	Q81kt9 plasmodium
43	104	9.3	1605	5	Q814X5	Q814x5 plasmodium
44	104	9.3	1859	5	Q81C27	Q81c27 plasmodium
45	104	9.3	1936	5	Q815A6	Q815a6 plasmodium

ALIGNMENTS

RESULT 1

Q8NUH0	PRELIMINARY;	PRT; 1183 AA.
ID	Q8NUH0	
AC	Q8NUH0;	
DT	01-OCT-2002 (Tremblrel. 22, Created)	
DT	01-OCT-2002 (Tremblrel. 22, Last sequence update)	
DT	01-OCT-2003 (Tremblrel. 25, Last annotation update)	
DE	Collagen adhesin.	
GN	MW2612.	
OS	Staphylococcus aureus (strain MW2).	
OC	Bacteria; Firmicutes; Bacillales; Staphylococcus.	
OX	NCBI_TaxID=196620;	
RI	[1]	
RP	SEQUENCE FROM N.A.	
RX	MEDLINE=22040717; PubMed=12044378;	
RA	Baba T., Takeuchi F., Kuroda M., Yuzawa H., Aoki K.-I., Oguchi A.,	
RA	Nagai Y., Iwana N., Asano K., Naimi T., Kuroda H., Cui L.,	
RA	Yamamoto K., Hiramatsu K.;	
RT	"Genome and virulence determinants of high virulence community-	
RT	acquired MRSA."	
RL	Lancet 359:1819-1827(2002).	
DR	EMBL; AF004831; BAB96477.1; -.	
DR	PIR; A42404; A42404.	
DR	PIR; A48620; A48620.	
DR	InterPro; IPR008966; Adhes_bact.	
DR	InterPro; IPR008454; Cna_B.	
DR	InterPro; IPR008970; Cna_B_unit.	
DR	InterPro; IPR008456; Collagen_bind.	
DR	InterPro; IPR001899; Gram_pos_anchor.	
DR	Pfam; PF05738; Cna_B_7.	
DR	Pfam; PF05737; Collagen_bind; 1.	
DR	TIGRFAM; TIGR01167; LPXG_anchor; 1.	
KW	Complete proteome.	
SQ	SEQUENCE 1183 AA; 133001 MW; 7A33DA876F9DA447 CRC64;	

Query Match 92.2%; Score 1035; DB 16; Length 1183;

Best Local Similarity 98.5%; Pred. No. 7.6e-58;

Matches 197; Conservative 1; Mismatches 2; Indels 0; Gaps 0;

```

QY 12 SDKVAITISGNKSTNVTVHKSEAGTSSVFFYKTDGMLPEDTTHVRWFLNINNEKSYVSK 71
DB 144 SDKVAITISGNKSTNVTVHKSEAGTSSVFFYKTDGMLPEDTTHVRWFLNINNEKRYVSK 203
QY 72 DITIKDQIQGGQDLSTLNINVTGTHSNYSQSAITDPEKAFPGSKITVDNKTNTIDV 131
DB 204 DITIKDQIQGGQDLSTLNINVTGTHSNYSQSAITDPEKAFPGSKITVDNKTNTIDV 263
QY 132 TIPOGYSNFSINYSKTIITNEQKEFVNNQAWYQEHGKEEVNGKSFNHTVHNINANA 191
DB 264 TIPOGYSNFSINYSKTIITNEQKEFVNNQAWYQEHGKEEVNGKSFNHTVHNINANA 323
QY 192 GIEGTVKGLKVLKQDKDK 211
DB 324 GIEGTVKGLKVLKQDKDK 343

RESULT 2
Q84BK5 PRELIMINARY; PRT; 721 AA.
AC Q84BK5;
DT 01-JUN-2003 (TrEMBLrel. 24, Created)
DT 01-JUN-2003 (TrEMBLrel. 24, Last sequence update)
DT 01-OCT-2003 (TrEMBLrel. 25, Last annotation update)
DE Collagen adhesin.
OS Enterococcus faecium (Streptococcus faecium).
OC Bacteria; Firmicutes; Lactobacillales; Enterococcaceae; Enterococcus.
OX NCBI_TaxID=1352;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=TX2555;
RX MEDLINE=22511562; PubMed=12622825;
RA Nallapareddy S.R., Weinstock G.M., Murray B.B.;
RT "Clinical isolates of Enterococcus faecium exhibit strain-specific collagen binding mediated by Acm, a new member of the MSCRAMM family.";
RL Mol. Microbiol. 47:1733-1747 (2003).
DR EMBL; AY135217; AAN12397.1; -.
DR InterPro; IPR008966; Adhes_bact.
DR InterPro; IPR008970; Cna_B_unit.
DR Pfam; PF00746; Gram_pos_anchor.
DR PROSITE; PS50847; GRAM_POS_ANCHORING; 1.
KW Collagen.
SQ SEQUENCE 721 AA; 81545 MW; DBSBB3B3DEF45673 CRC64;

Query Match 43.8%; Score 491; DB 2; Length 721;
Best Local Similarity 48.5%; Pred. No. 2e-23;
Matches 95; Conservative 35; Mismatches 64; Indels 2; Gaps 2;

QY 18 TITSGNKSNTVTVHKSEAG-TSSVFFYKTDGMLPEDTTHVRWFLNINNEKSYVSKDITIK 76
DB 150 TVTSGDKTATVNTKPSAGSSSSVFFYKTDGMLPEDTTHVRWFLNINNGTYVEQPVKIS 209
QY 77 DOIQGGQDLSTLNIN-VTGTHSNYSQSAITDPEKAFPGSKITVDNKTNTIDVTIPQ 135
DB 210 DEIQSGQRDPSTFELNQLHLEQKVRGEGIQQLQFPSPATFNSVDNYIETIPK 269
QY 136 GYGSYNSFSINYSKTIITNEQKEFVNNQAWYQEHGKEEVNGKSFNHTVHNINANAGIEG 195
DB 270 NFVNLRLKIMVSYKTIENPEQINFENHSEAWFKNKPAVDGESFNHTVKNISASGGVNG 329
QY 196 TVKGLKVLKQDKDK 211
DB 330 TVRGLKIPKIFYNDTE 345

RESULT 3
Q84OA5 PRELIMINARY; PRT; 657 AA.
AC Q84OA5;
DT 01-JUN-2003 (TrEMBLrel. 24, Created)
DT 01-JUN-2003 (TrEMBLrel. 24, Last sequence update)

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DT 01-OCT-2003 (TrEMBLrel. 25, Last annotation update)
DE Collagen-binding protein Cne precursor (Fragment).
GN CNE.
OS Streptococcus equi subsp. equi.
OC Bacteria; Firmicutes; Lactobacillales; Streptococcaceae;
OC Streptococcus.
OX NCBI_TaxID=148942;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=1866;
RA Lannergard J., Frykberg L., Guss B.;
RT "CNE, a collagen-binding protein of Streptococcus equi.";
RL Submitted (DSC-2002) to the EMBL/GenBank/DBJ databases.
DR EMBL; AY193773; AAO43099.1; -.
DR InterPro; IPR008966; Adhes_bact.
DR InterPro; IPR008970; Cna_B_unit.
DR InterPro; IPR001899; Gram_pos_anchor.
DR TIGRFAMs; TIGR01167; LPXTG_anchor; 1.
KW Collagen; Signal.
FT SIGNAL 1 27 Potential.
FT NON_TER 657
SQ SEQUENCE 657 AA; 73889 MW; A902B85BBCE59C4F CRC64;

Query Match 41.3%; Score 463.5; DB 2; Length 657;
Best Local Similarity 47.7%; Pred. No. 1e-21;
Matches 94; Conservative 36; Mismatches 62; Indels 5; Gaps 3;

QY 18 TITSGNKSNTVTVHKSEAGT-SVFFYKTDGMLPEDTTHVRWFLNINNEKSYVSKDITIK 76
DB 146 TVKVGNRATITVTKPEAGTGTSSVFFYKTDGMDPNDTERTVWFLNINNEKSWANTVTVE 205
QY 77 DOIQGGQDLSTLNINVTGTHSNYSQSAITDPEKAFPGSKITVDNKTNTIDVTIPQ 136
DB 206 DDIOGGQDLMSFDITVSGYRNERFVGENALTFFHTTTPNSVITA--TDNHSIVRLDQY 263
QY 137 YGSYNSFSINYSKTIITNEQKEFVNNQAWYQEHGKEEVNGKSFNHTVHNINANAGIEGT 196
DB 264 DASQNTVNIAYKTIITDQKEFANNKIKWQILYKQVSGQESNHNQANINANGVDGS 323
QY 197 --VRGELKVLKQDKDK 211
DB 324 RYTSFTVKKIWNDEKQ 340

RESULT 4
Q8KRZ5 PRELIMINARY; PRT; 221 AA.
AC Q8KRZ5;
DT 01-OCT-2002 (TrEMBLrel. 22, Created)
DT 01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
DE Putative collagen adhesin se73.9 (Fragment).
OS Streptococcus equi.
OC Bacteria; Firmicutes; Lactobacillales; Streptococcaceae;
OC Streptococcus.
OX NCBI_TaxID=1336;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=CF32;
RA Qin A., Artushin S.C., Timoney J.F.;
RT "Identification and Genomic Organization of Genes for Immunoreactive Surface Exposed Proteins of Streptococcus equi.";
RL Submitted (JUN-2002) to the EMBL/GenBank/DBJ databases.
DR EMBL; AF521599; AAM77657.1; -.
DR InterPro; IPR008966; Adhes_bact.
DR InterPro; IPR008970; Cna_B_unit.
DR InterPro; IPR008456; Collagen_bind.
DR Pfam; PF05737; Collagen_bind; 1.
KW Collagen.
FT NON_TER 1
FT NON_TER 221
SQ SEQUENCE 221 AA; 25049 MW; 0300F430D21EB859 CRC64;

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Query Match 41.2%; Score 462.5; DB 2; Length 221;
Best Local Similarity 47.7%; Pred. No. 3.4e-22;
Matches 94; Conservative 36; Mismatches 62; Indels 5; Gaps 3;

QY 18 TITSGNKSTNVTVHKSEAGT-SSVFYKGTGDMLPEDTTHVRWFLNINNEKSYVSKDITIK 76
DB 2 TVKVGRTCTITVKEAGTGTSSFYKGTGDMQPNDETRVRWFLNINNEKSYVSKDITIK 61
QY 77 DQIQGGQQLDLSTLNLINVTGTHSNYSYGOSAITDFEKAFFGSKITVDNTKNTIDVTIPQ 136
DB 62 DDQIQGGQQLDLSTLNLINVTGTHSNYSYGOSAITDFEKAFFGSKITVDNTKNTIDVTIPQ 119
QY 137 YGSYNSFSNYTKITNEQKQFVNNSQAWYQDHGKEVNGKSFNHTVHNINAGIEGT 196
DB 120 DASQNTVNIAYTKITDFDQKEFANNSKIWIYQLYKQVSGQSNHQVANNANGVDGS 179
QY 197 --VKGELKVLKQDKTK 211
DB 180 RYTSFTVKKIWNDEKQ 196

RESULT 5
Q7WMP3 PRELIMINARY; PRT; 1151 AA.
AC Q7WMP3;
DT 01-OCT-2003 (TReMBLrel. 25, Created)
DT 01-OCT-2003 (TReMBLrel. 25, Last sequence update)
DE Collagen-binding protein A precursor.
GN CBPA.
OS Arcanobacterium pyogenes.
OC Bacteria; Actinobacteria; Actinobacteridae; Actinomycetales;
OC Actinomycineae; Actinomycetaceae; Arcanobacterium.
OX NCBI_TaxID=1661;
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE=22756333; PubMed=12874314;
RA Esmy P.A., Billington S.J., Link M.A., Songer J.G., Jost B.H.;
RT "The Arcanobacterium pyogenes Collagen-Binding Protein, Cbpa, Promotes
RT Adhesion to Host Cells."
RL Infect. Immun. 71:4368-4374(2003).
DR EMBL; AY223543; AAO43108.1; -.
KW Collagen; Signal.
FT SIGNAL 1 27 Potential.
SQ SEQUENCE 1151 AA; 124719 MW; 3127352B9795CE98 CRC64;

Query Match 16.3%; Score 182.5; DB 2; Length 1151;
Best Local Similarity 29.3%; Pred. No. 0.0016;
Matches 61; Conservative 34; Mismatches 91; Indels 21; Gaps 8;

QY 18 TITSGNKSTNVTVHKSEAGTSSVFYKGTGDMLPEDTT-HVRWFLNIN-NEKSYVSKDITI 75
DB 156 TVVGN--TAVQISHNRGDRGVFAGTKGLYGSNPGYVRWFLRANINGDPWPGGDVI 213
QY 76 KDQIQGGQQLDLSTLNLINVTGTHSN-----YVSGOSAITDFEKAFFGSK-----ITVDNT 125
DB 214 EDTLIGIQKLDGTGIRI---GLHWGQRPGSYSLTWSIEDFLNSDYGTRAGTIAYKEH 270
QY 126 KNTIDVTIPQYGSYNSFSNYTKITNEQKQFVNNSQ-AWYQDHGKEVNGKSFNHTV 184
DB 271 EGRISHVPOSVVNGREFSFTYDAKIMDDTQBEFTNTATDFYENQKORIVD---IYTV 327
QY 185 HNNANAGIEGTVKGELKVLKQDKTK 211
DB 328 RNPASGGIEGKTTASLNLDKVLKDSR 354

RESULT 6
Q9F859 PRELIMINARY; PRT; 319 AA.
AC Q9F859;
DT 01-MAR-2001 (TReMBLrel. 16, Created)
DT 01-MAR-2001 (TReMBLrel. 16, Last sequence update)
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DT 01-OCT-2003 (TReMBLrel. 25, Last annotation update)
DE Collagen adhesin (Fragment).
GN ACE.
OS Enterococcus faecalis (Streptococcus faecalis).
OC Bacteria; Firmicutes; Lactobacillales; Enterococcaceae; Enterococcus.
OX NCBI_TaxID=1351;
RN [1]
RP SEQUENCE FROM N.A.
RX STRAIN=Beirut/TX0645;
RC MEDLINE=20407335; PubMed=10948146;
RA Nallapareddy S.R., Singh K.V., Duh R.W., Weinstock G.M., Murray B.E.;
RT "Diversity of ace, a gene encoding a microbial surface component
RT recognizing adhesive matrix molecules, from different strains of
RT enterococcus faecalis and evidence for production of ace during human
RT infections."
RL Infect. Immun. 68:5210-5217(2000).
DR EMBL; AF260889; AAG23947.1; -.
DR HSSP; Q53654; 1AMX.
DR InterPro; IPR008966; Adhes_bact.
DR InterPro; IPR008456; Collagen_bind.
DR Pfam; PF05737; Collagen_bind; 1.
FT NON_TER 1 319
FT NON_TER 319 319
SQ SEQUENCE 319 AA; 35986 MW; D5F914E0856A6940 CRC64;

Query Match 16.2%; Score 182; DB 2; Length 319;
Best Local Similarity 28.8%; Pred. No. 0.00039;
Matches 55; Conservative 32; Mismatches 88; Indels 16; Gaps 6;

QY 18 TITSGNKSTNVTVHKSEAG---TSSVFYKGTGDMLPEDTTHVRWFLNINNEKSYVSKDIT 74
DB 112 TATATQRLTIEGVTNTETGTQIERDYFFYKVGDLAGE-SNQVRWFLNVLNKSVDVTIS 170
QY 75 IKDQIQGGQQLDLSTLNLINVTGTHSNYSYGOSAITDFEKAFFGSKITVDNTKNTIDVTIP 134
DB 171 IADROGSGQQLNKESFTFDIVNDKETY---ISLAEFEQGGYKIDFV--TNDNFNLFY 225
QY 135 QYGSYNSFSNYTKIT--NEQKQFVNNSQAWYQDHGKEVNGKSFNHTVHNINAG 192
DB 226 QDKARFTSFIVRYTSTITEAGQHQATFENSVDINTYQLNNQDATNEKTSQV-----KNVF 280
QY 193 IEGTVKGLKV 203
DB 281 VEGEASGNQNV 291

RESULT 7
Q9F866 PRELIMINARY; PRT; 721 AA.
AC Q9F866;
DT 01-MAR-2001 (TReMBLrel. 16, Created)
DT 01-MAR-2001 (TReMBLrel. 16, Last sequence update)
DT 01-OCT-2003 (TReMBLrel. 25, Last annotation update)
DE Collagen adhesin precursor.
GN ACE.
OS Enterococcus faecalis (Streptococcus faecalis).
OC Bacteria; Firmicutes; Lactobacillales; Enterococcaceae; Enterococcus.
OX NCBI_TaxID=1351;
RN [1]
RP SEQUENCE FROM N.A.
RX STRAIN=END6/TX0045;
RC MEDLINE=20407335; PubMed=10948146;
RA Nallapareddy S.R., Singh K.V., Duh R.W., Weinstock G.M., Murray B.E.;
RT "Diversity of ace, a gene encoding a microbial surface component
RT recognizing adhesive matrix molecules, from different strains of
RT enterococcus faecalis and evidence for production of ace during human
RT infections."
RL Infect. Immun. 68:5210-5217(2000).
DR EMBL; AF260873; AAG23932.1; -.
DR HSSP; Q53654; 1AMX.
DR InterPro; IPR008966; Adhes_bact.
DR InterPro; IPR008456; Collagen_bind.
DR InterPro; IPR001899; Gram_pos_anchor.
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QY 135 QYGSYNSPISYNYTKIT--NEOQKEFVNNSQAWYQEHGKEVNGKSFNHTVHNINANAG 192
Db 226 RDKARFTSFIVRYTSTITEAGHQATFENSIDYINQLNNQDATNEKNTSQV-----KNVF 280
QY 193 IEGTVKGELKV 203
Db 281 VEGEASGNQNV 291

RESULT 13
QXQBQ7
ID Q9XBQ7 PRELIMINARY; PRT; 458 AA.
AC Q9XBQ7;
DT 01-NOV-1999 (TREMBlrel. 12, Created)
DT 01-NOV-1999 (TREMBlrel. 12, Last sequence update)
DT 01-OCT-2003 (TREMBlrel. 25, Last annotation update)
DE Collagen adhesin Ace (Fragment).
GN ACE.
OS Enterococcus faecalis (Streptococcus faecalis).
OC Bacteria; Firmicutes; Lactobacillales; Enterococcaceae; Enterococcus.
OX NCBI_TaxID=1351;
RN [1]
RP SEQUENCE FROM N.A.
RX STRAIN=CG110;
RX MEDLINE=99410431; PubMed=10480905;
RA Rich R.L., Kreikeneyer B., Owens R.T., LaBrenz S., Narayana S.V.,
RA Weinstock G.M., Murray B.E., Hook M.;
RT "Ace is a collagen-binding MSCRAMM from Enterococcus faecalis.";
RL J. Biol. Chem. 274:26939-26945(1999).
DR EMBL; AF159247; AAD43342.1; -.
DR HSSP; Q53654; 1AMX.
DR InterPro; IPR008966; Adhes_bact.
DR InterPro; IPR008456; Collagen_bind.
DR Pfam; PF05737; Collagen_bind; 1.
FT NON_TER 1
FT NON_TER 458 458
SQ SEQUENCE 458 AA; 51085 MW; B674692BD29A0A24 CRC64;

Query Match 15.9%; Score 178; DB 2; Length 458;
Best Local Similarity 28.3%; Pred. No. 0.0011;
Matches 54; Conservative 33; Mismatches 88; Indels 16; Gaps 6;

QY 18 TITSGNKSTNVTVHKSEAG---TSSVFYKGTGMLPETHVRWFLNINNEKSYVKDIT 74
Db 121 TATATQRLTIEGVTNTETQIERDYPFFYKVGDLAGE-SNQVRWFLNVLNKSVDTEIS 179
QY 75 IKDQIOGGQQLDLSTLINVGTGTHSNYSGSAITDFEKAFFGSKITVDNTKNTIDVTIP 134
Db 180 IADRQSGGQQLNKESFTFDIVNDKETKY---ISLAEFEQGGYKIDFV--TDNDFNLRFY 234
QY 135 QYGSYNSPISYNYTKIT--NEOQKEFVNNSQAWYQEHGKEVNGKSFNHTVHNINANAG 192
Db 235 RDKARFTSFIVRYTSTITEAGHQATFENSIDYINQLNNQDATNEKNTSQV-----KNVF 289
QY 193 IEGTVKGELKV 203
Db 290 VEGEASGNQNV 300

RESULT 14
Q9F862
ID Q9F862 PRELIMINARY; PRT; 580 AA.
AC Q9F862;
DT 01-MAR-2001 (TREMBlrel. 16, Created)
DT 01-MAR-2001 (TREMBlrel. 16, Last sequence update)
DT 01-OCT-2003 (TREMBlrel. 25, Last annotation update)
DE Collagen adhesin precursor.
GN ACE.
OS Enterococcus faecalis (Streptococcus faecalis).
OC Bacteria; Firmicutes; Lactobacillales; Enterococcaceae; Enterococcus.
OX NCBI_TaxID=1351;
RN [1]
RP SEQUENCE FROM N.A.
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RC STRAIN=LBJ-1/TX0020;
RX MEDLINE=20407335; PubMed=10948146;
RA Nallapareddy S.R., Singh K.V., Duh R.W., Weinstock G.M., Murray B.E.;
RT "Diversity of ace, a gene encoding a microbial surface component
RT recognizing adhesive matrix molecules, from different strains of
RT enterococcus faecalis and evidence for production of ace during human
RT infections.";
RL Infect. Immun. 68:5210-5217(2000).
DR EMBL; AF260878; AAG233936.1; -.
DR HSSP; Q53654; 1AMX.
DR InterPro; IPR008966; Adhes_bact.
DR InterPro; IPR008456; Collagen_bind.
DR InterPro; IPR001899; Gram_pos_anchor.
DR Pfam; PF05737; Collagen_bind; 1.
DR Pfam; PF00746; Gram_pos_anchor; 1.
DR PROSITE; PS50847; GRAM_POS_ANCHORING; 1.
KW Signal.
FT SIGNAL 1 31
SQ SEQUENCE 580 AA; 63948 MW; 7CD61EAD3FDA0993 CRC64;

Query Match 15.9%; Score 178; DB 2; Length 580;
Best Local Similarity 28.3%; Pred. No. 0.0014;
Matches 54; Conservative 33; Mismatches 88; Indels 16; Gaps 6;

QY 18 TITSGNKSTNVTVHKSEAG---TSSVFYKGTGMLPETHVRWFLNINNEKSYVKDIT 74
Db 152 TATATQRLTIEGVTNTETQIERDYPFFYKVGDLAGE-SNQVRWFLNVLNKSVDTEIS 210
QY 75 IKDQIOGGQQLDLSTLINVGTGTHSNYSGSAITDFEKAFFGSKITVDNTKNTIDVTIP 134
Db 211 IADRQSGGQQLNKESFTFDIVNDKETKY---ISLAEFEQGGYKIDFV--TDNDFNLRFY 265
QY 135 QYGSYNSPISYNYTKIT--NEOQKEFVNNSQAWYQEHGKEVNGKSFNHTVHNINANAG 192
Db 266 RDKARFTSFIVRYTSTITEAGHQATFENSIDYINQLNNQDATNEKNTSQV-----KNVF 320
QY 193 IEGTVKGELKV 203
Db 321 VEGEASGNQNV 331

RESULT 15
Q9F861
ID Q9F861 PRELIMINARY; PRT; 580 AA.
AC Q9F861;
DT 01-MAR-2001 (TREMBlrel. 16, Created)
DT 01-MAR-2001 (TREMBlrel. 16, Last sequence update)
DT 01-OCT-2003 (TREMBlrel. 25, Last annotation update)
DE Collagen adhesin precursor.
GN ACE.
OS Enterococcus faecalis (Streptococcus faecalis).
OC Bacteria; Firmicutes; Lactobacillales; Enterococcaceae; Enterococcus.
OX NCBI_TaxID=1351;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=JH2-2;
RX MEDLINE=20407335; PubMed=10948146;
RA Nallapareddy S.R., Singh K.V., Duh R.W., Weinstock G.M., Murray B.E.;
RT "Diversity of ace, a gene encoding a microbial surface component
RT recognizing adhesive matrix molecules, from different strains of
RT enterococcus faecalis and evidence for production of ace during human
RT infections.";
RL Infect. Immun. 68:5210-5217(2000).
DR EMBL; AF260879; AAG233937.1; -.
DR HSSP; Q53654; 1AMX.
DR InterPro; IPR008966; Adhes_bact.
DR InterPro; IPR008456; Collagen_bind.
DR InterPro; IPR001899; Gram_pos_anchor.
DR Pfam; PF05737; Collagen_bind; 1.
DR Pfam; PF00746; Gram_pos_anchor; 1.
DR PROSITE; PS50847; GRAM_POS_ANCHORING; 1.
KW Signal.
FT SIGNAL 1 31
SQ SEQUENCE 580 AA; 63948 MW; 7CD61EAD3FDA0993 CRC64;
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SQ SEQUENCE 580 AA; 64176 MW; 060EFES0D61A5271 CRC64;  
Query Match 15.9%; Score 178; DB 2; Length 580;  
Best Local Similarity 28.3%; Pred. No. 0.0014;  
Matches 54; Conservative 33; Mismatches 88; Indels 16; Gaps 6;  
QY 18 TITSGNKSTNVTVHKSEAG---TSSVFYKTDMLPEDTTHVRWFLNINNEKSYVSKDIT 74  
DB 152 TATATORLTIEGVNTETGQIERDYPFFYKVGDLAGE-SNQVRWFLNVLNLSKSDVTEDIS 210  
QY 75 IKDQIQGGQQLDLSTLNIINVTGTHSNVYGSQSAITDFEKAFPGSKITVDNTKNTIDVTIP 134  
DB 211 IADRQSGGQQLNKESFTFDIVNDKETY---ISLAEFEQOGYKIDFV--TDNDFNLRFY 265  
QY 135 QGYGSYNFSINYYTKKIT--NEQQKEFVANSQAWYQEHGKEEVNGKSFNHTVHNINANAG 192  
DB 266 RDKARFTSFIVRYTSTITEAGQHQATFENSYDINYQLNNQDATNEKNTSQV-----KNVF 320  
QY 193 IEGTVKGELKV 203  
DB 321 VEGEASGNQNV 331

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Job time : 42 secs

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